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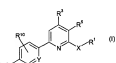
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(54) Title: COX-2 INHIBITING PYRIDINE DERIVATIVES

16970/7007 O R¹O₂s



(57) Abstract: Compounds of formula (I) or pharmaceutically acceptable salts thereof are potent and selective inhibitors of COX-2 and are of use in the treatment of the pain, fever and inflammation of a variety of conditions and diseases.

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COX-2 INHIBITING PYRIDINE DERIVATIVES

This invention relates to pyridine derivatives, to processes for their preparation, to pharmaceutical compositions containing them and to their use in medicine.

The enzyme cyclooxygenase (COX) has recently been discovered to exist in two isoforms, COX-1 and COX-2. COX-1 corresponds to the originally identified constitutive enzyme while COX-2 is rapidly and readily inducible by a number of agents including mitogens, endotoxin, hormones, cytokines and growth factors. Prostaglandins generated by the action of COX have both physiological rolations at it is generally believed that COX-1 is largely responsible for the important physiological functions such as maintenance of gastrointestinal integrity and renal blood flow. In contrast the inducible form, COX-2, is believed to be largely responsible for the pathological effects of prostaglandins where rapid induction of the enzyme occurs in response to such agents as inflammatory agents, hormones, growth factors and cytokines. A selective inhibitor of COX-2 would therefore have anti-inflammatory, anti-pyretic and analgesic properties, without the potential side effects associated with inhibition of COX-1. We have now found a novel group of compounds which are both potent and selective inhibitors of COX-2.

The invention thus provides a compound of formula (I)

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(1)

or a pharmaceutically acceptable salt thereof in which:

X is selected from the group consisting of oxygen or NR²;
Y is selected from the group consisting of CH or nitrogen;
R¹ is selected from the group consisting of H, C₁₋₈alkyl, C₁₋₂alkyl substituted by one to five fluorine atoms, C₁₋₃alkylOC₁₋₃alkyl, C₃₋₆alkenyl, C₃₋₆alkynyl, C₃₋₁₀cycloalkylC₀₋₆alkyl, C₄₋₇cycloalkyl substituted by C₁₋₃alkyl or C₁₋₃alkoxy, C₄₋₇bridoed cycloalkyl, A(CR[®]R⁷)_n and B(CR[®]R⁷)_n:

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R2 is selected from the group consisting of H and C1-6alkyl; or

R¹ and R², together with the nitrogen atom to which they are attached form a 4-8 membered saturated heterocyclic ring such as a pyrrolidine, morpholine or piperidine ring, or a 5-membered heteroaryl ring which is unsubstituted or substituted by one R⁸:

 R^3 is selected from the group consisting of C_{1-5} alkyl and C_{1-2} alkyl substituted by one to five fluorine atoms;

R⁴ is selected from the group consisting of C₁₋₆alkyl, NH₂ and R⁹CONH;

 R^5 is selected from the group consisting of hydrogen, C_{1-3} alkyl, C_{1-2} alkyl substituted by one to five fluorine atoms, C_{1-3} alkyl O_2 C, halogen, cyano, $(C_{1-3}$ alkyl $)_2$ NCO, C_{1-3} alkyl $)_3$ NCO, C_{1-3} alkyl $)_4$ NCO, C_{1-3} alkyl $)_5$ NCO, $C_$

R⁶ and R⁷ are independently selected from H or C₁₋₆alkyl;

A is an unsubstituted 5- or 6-membered heteroaryl or an unsubstituted 6membered aryl, or a 5- or 6-membered heteroaryl or a 6-membered aryl substituted by one or more R⁸:

 R^8 is selected from the group consisting of halogen, C_{1-6} alkyl, C_{1-6} alkyl substituted by one more fluorine atoms, C_{1-6} alkoxy, C_{1-6} alkoxy substituted by one or more F, NH_2SO_2 and C_{1-6} alkyl SO_2 ;

B is selected from the group consisting of

20 defines the point of attachment of the ring;

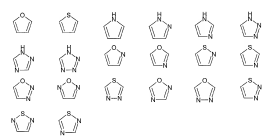
 R^9 is selected from the group consisting of H, $C_{1:6}$ alkyl, $C_{1:6}$ alkyo, $C_{1:6}$ alkyl, phenyl, HO $_2$ CC $_{1:6}$ alkyl, $C_{1:6}$ alkyl, COCO $_{1:6}$ alkyl, C $_{1:6}$

 R^{10} is selected from the group consisting of H and halogen; and n is 0 to 4.

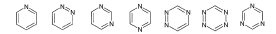
The term 'halogen' is used to represent fluorine, chlorine, bromine or iodine.

The term 'alkyl' as a group or part of a group means a straight or branched chain alkyl group, for example a methyl, ethyl, n-propyl, i-propyl, n-butyl, s-butyl or t-butyl group.

30 The term 'saturated heterocyclic' means a saturated ring containing at least one atom other than carbon. The term '5-membered heteroaryl' means a heteroaryl selected from the following:



5 The term '6- membered heteroaryl' means a heteroaryl selected from the following:



The term '6-membered aryl' means:



It is to be understood that the present invention encompasses all isomers of the compounds of formula (I) and their pharmaceutically acceptable derivatives, including all geometric, tautomeric and optical forms, and mixtures thereof (e.g. racemic mixtures). In particular when the ring B lacks a plane of symmetry the compounds of formula (I) contain a chiral centre as indicated therein by the asterisk *. Furthermore, it will be appreciated by those skilled in the art that when R⁶ and R⁷ in formula (I) are different the corresponding compounds contain at least one chiral centre, by virtue of the asymmetric carbon atom defined thereby,

and that such compounds exist in the form of a pair of optical isomers (i.e. enantiomers).

It will be appreciated that in some instances, compounds of the present invention may include a basic function such as an amino group as a substituent. Such basic functions may be used to form acid addition salts, in particular pharmaceutically acceptable salts. Pharmaceutically acceptable salts include those described by Berge, Bighley, and Monkhouse, J. Pharm. Sci., 1977, 66, 1-19. Such salts may be formed from inorganic and organic acids. Representative examples thereof include maleic, fumaric, benzoic, ascorbic, pamoic, succinic, bismethylenesalicylic, methanesulfonic, ethanedisulfonic, acetic, propionic, tartaric, salicylic, citric, gluconic, aspartic, stearic, palmitic, itaconic, glycolic, p-aminobenzoic, glutamic, taurocholic acid, benzenesulfonic, phosphoric and nitric acids.

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It will be appreciated that in some instances, compounds of the present invention may include a carboxy group as a substituent. Such carboxy groups may be used to form salts, in particular pharmaceutically acceptable salts. Pharmaceutically acceptable salts include those described by Berge, Bighley, and Monkhouse, *J. Pharm. Sci.*, 1977, **66**, 1-19. Preferred salts include alkali metal salts such as the sodium and potassium salts.

In one aspect the invention provides a compound of formula (IA)

(IA)

or a pharmaceutically acceptable salt thereof in which:

X is selected from the group consisting of oxygen or NR²; Y is selected from the group consisting of CH or nitrogen; R^1 is selected from the group consisting of H, $C_{1\text{--}8}$ alkyl, $C_{1\text{--}2}$ alkyl substituted by one to five fluorine atoms, $C_{1\text{--}3}$ alkyl, $C_{1\text{--}3}$ alkyl, $C_{3\text{--}8}$ alkenyl, $C_{3\text{--}8}$ alkynyl, $C_{3\text{--}8}$ alkyl, $C_{4\text{--}12}$ bridged cycloalkyl, $A(CR^6R^7)_n$ and $B(CR^6R^7)_n$;

R2 is selected from the group consisting of H and C1-6alkyl; or

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R¹ and R², together with the nitrogen atom to which they are attached form a 4-8 membered saturated heterocyclic ring such as a pyrrolidine, morpholine or piperidine ring;

 R^3 is selected from the group consisting of C₁₋₅alkyl and C₁₋₂alkyl substituted by one to five fluorine atoms:

R⁴ is selected from the group consisting of C_{1.6}alkyl, NH₂ and R⁹CONH;

 R^5 is selected from the group consisting of hydrogen, C_{1-3} alkyl, C_{1-2} alkyl substituted by one to five fluorine atoms, halogen, cyano, $(C_{1-3}$ alkyl)₂NCO, C_{1-3} alkyl and C_{1-3} alkylO₂S:

R⁶ and R⁷ are independently selected from H or C₁₋₆alkyl;

A is an unsubstituted 5- or 6-membered heteroaryl or an unsubstituted 6-membered aryl, or a 5- or 6-membered heteroaryl or a 6-membered aryl substituted by one or more R⁸:

R⁸ is selected from the group consisting of halogen, C₁₋₆alkyl, C₁₋₆alkyl substituted by one more fluorine atoms, C₁₋₆alkoxy, C₁₋₆alkoxy substituted by one or more F. NH₂SO₂ and C₁₋₆alkylSO₂;

B is selected from the group consisting of

defines the point of attachment of the ring;

 R^9 is selected from the group consisting of H, C_{1-6} alkyl, C_{1-6} alkyl, C_{1-6} alkyl, phenyl, HO_2CC_{1-6} alkyl, C_{1-6} alkyl

 R^{10} is selected from the group consisting of H and halogen; and n is 0 to 4.

In another aspect the invention provides a compound of formula (IB)

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$$R^{4}O_{2}S$$
 $P^{1}O_{3}S$
 $P^{1}O_{4}S$
 $P^{1}O_{5}S$
 $P^{1}O_{5}S$

or a pharmaceutically acceptable salt thereof, in which all substituents are as for a compound of formula (I) defined hereinabove.

In another aspect the invention provides a compound of formula (IC)

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or a pharmaceutically acceptable salt thereof in which:

X is selected from the group consisting of oxygen or NR²;

Y is selected from the group consisting of CH or nitrogen;

 R^{1} is selected from the group consisting of H, $C_{1:\delta}$ alkyl, $C_{1:2}$ alkyl substituted by one to five fluorine atoms, $C_{1:3}$ alkyl, $C_{1:3}$ alkyl, $C_{3:\delta}$ alkenyl, $C_{3:\delta}$ alkynyl, $C_{3:\delta}$ $_{10}$ cycloalkyl, $C_{4:7}$ cycloalkyl substituted by $C_{1:3}$ alkyl or $C_{1:3}$ alkoxy, $C_{4:7}$ bridged cycloalkyl, $A(CR^{6}R^{7})_{n}$ and $B(CR^{6}R^{7})_{n}$;

R2 is selected from the group consisting of H and C1-6alkyl; or

R¹ and R², together with the nitrogen atom to which they are attached form a 4-8 membered saturated heterocyclic ring such as a pyrrolidine, morpholine or piperidine ring, or a 5-membered heteroaryl ring which is unsubstituted or substituted by one R⁸.

 R^3 is selected from the group consisting of $\mathsf{C}_{1.5}$ alkyl and $\mathsf{C}_{1.2}$ alkyl substituted by one to five fluorine atoms:

R⁴ is selected from the group consisting of C₁₋₈alkyl, NH₂ and R⁹CONH; R⁵ is selected from the group consisting of hydrogen, C₁₋₃alkyl, C₁₋₂alkyl substituted by one to five fluorine atoms, C₁₋₃alkylO₂C, halogen, cyano, (C₁₋₃alkyl)₂NCO, C₁₋₃alkylS and C₁₋₃alkylO₂S;

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R⁶ and R⁷ are independently selected from H or C₁₋₆alkyl;

A is an unsubstituted 5- or 6-membered heteroaryl or an unsubstituted 6-membered aryl, or a 5- or 6-membered heteroaryl or a 6-membered aryl substituted by one or more R⁸:

R⁸ is selected from the group consisting of halogen, C₁₋₈alkyl, C₁₋₈alkyl substituted by one more fluorine atoms, C₁₋₈alkoxy, C₁₋₈alkoxy substituted by one or more F, NH₂SO₂ and C₁₋₈alkylSO₂;

B is selected from the group consisting of

$$\rightarrow \bigcirc$$
 , $\rightarrow \bigcirc$, $\rightarrow \bigcirc$, $\rightarrow \bigcirc$ and where

defines the point of attachment of the ring;

 R^{9} is selected from the group consisting of H, C_{1-6} alkyl, C_{1-6} alkyl, C_{1-6} alkyl, phenyl, $HO_{2}CC_{1-6}$ alkyl, C_{1-6} a

15 In another aspect of the invention Y is carbon.

In another aspect of the invention R^1 is selected from the group consisting of, $C_{1-galkyl}$, C_{3-10} cycloalkyl $C_{0-galkyl}$, C_{5-6} cycloalkyl substituted by $C_{1-2alkyl}$ or $C_{1-2alkyl}$ or $C_{1-2alkyl}$ and $C_{1-2alkyl}$ substituted by one to five fluorine atoms.

Representative examples of R¹ include cyclohexylmethyl, cyclohexyl, n-butyl, npentyl, cyclopentyl, 2-methylpropyl, 2,2-dimethylpropyl, 2,2,2-trifluoroethyl, 2methoxyethyl and ethyl.

Further representative examples of R¹ include 1-methylethyl, 1-ethylpropyl, cycloheptyl, cis-4-methylcyclohexyl, trans-4-methylcyclohexyl, cyclobutyl, cyclopentanemethyl, and trans-4-(ethoxy)cyclohexyl.

In another aspect of the invention R^1 is selected from the group consisting of $A(CR^6R^7)_n$ and $B(CR^6R^7)_n$.

Further representative examples of R¹ include benzyl, 4-chlorobenzyl, 2-furylmethyl, 4-methylphenyl, 4-fluorophenyl, 4-methoxyphenyl, 3-pyridyl, 2-

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chlorophenyl, 3,5-difluorobenzyl, 3-pyridylmethyl, 2-methylbenzyl, 2-chlorobenzyl, (S)- α -methylbenzyl, (R)- α -methylbenzyl, 6-methylpyridin-3-yl, 4-methybenzyl, 4-fluorobenzyl, 2-(5-methylfuyrl)methyl, 4-methylbenzyl, 4-pyridylmethyl, 2-pyridylmethyl, 2-(6-methylpyridine)methyl, 2-thiophenylmethyl, 4-pyranylmethyl, 2-tetrahydrofurylmethyl, 2-(5-methylpyrazine)methyl and 4-ethoxybenzyl.

Further representative examples of R1 include 1H-imidazol-2-vlmethvl. 1Hpyrazol-4-vlmethyl, (1-methyl-1H-imidazol-2-yl)methyl, (3-methyl-1H-pyrazol-4vI)methyl. (1-methyl-1H-pyrazol-3-vI)methyl. (1-methyl-1H-pyrazol-4-vI)methyl. (3-methyl-1H-pyrazol-5-yl)methyl, (1-methyl-1H-pyrazol-5-yl)methyl, (1-methyl-1H-1,2,4-triazol-5-yl)methyl, (5-methyl-3-isoxazolyl)methyl, tetrahydro-2H-pyrantetrahydro-2H-pyran-4-ylmethyl, (6-methyl-3-pyridyl)methyl, pyrazinylmethyl, (2-methyl-1H-imidazol-4-yl)methyl, (4-methyl-1H-imidazol-5vI)methyl, (4-methyl-1H-imidazol-2-yI)methyl, (1-ethyl-1H-imidazol-2-yI)methyl, (1,3-dimethyl-1H-pyrazol-4-yl)methyl, (1,5-dimethyl-1H-pyrazol-4-yl)methyl, (3methyl-5-isothiazolyl)methyl, (4-methyl-1,3-thiazol-2-yl)methyl, (3-methyl-4-[1-(fluoromethyl)-1H-pyrazol-4-yl]methyl, (2-methyl-3isothiazolyl)methyl, pyridyl)methyl, (6-methyl-3-pyridyl)methyl, (1-methyl-1H-imidazol-2-yl)methyl. (5chloro-2-pyridyl)methyl, 1H-imidazol-2-ylmethyl, 4-ethoxyphenyl, 3-chloro-4methylphenyl, (5-chloro-2-pyridyl)methyl, (6-methyl-3-pyridyl)methyl, 2-methyl-3pyridyl, 6-methyl-2-pyridyl, 2-pyrazinylmethyl, 2,6-dimethyl-3-pyridyl, 3,4dichlorobenzyl, 5-chloro-3-pyridyl, 6-chloro-3-pyridazinyl, 3,5-dichlorobenzyl, 2carboxyphenyl, (5-methyl-2-pyridyl)methyl, 4-chloro-3-(trifluoromethyl)benzyl, (5bromo-2-pyridyl)methyl, (4-bromo-4-pyridyl)methyl, isoxazolyl)methyl, 5-pyrimidinylmethyl, (3-methyl-1,2,4-oxadiazol-5-yl)methyl, (5-

In another aspect of the invention R¹ is selected from the group consisting of C₃. salkenyl and C₃.salkynyl.

methyl-1,2,4-oxadiazol-3-yl)methyl and (1-ethyl-1H-1,2,4-triazol-5-yl)methyl.

Further representative examples of R¹ include propargyl and allyl.

30 In another aspect of the invention R² is H or C₁₋₂alkyl.

Representative examples of R2 include H, methyl and ethyl.

In another aspect of the invention \mbox{R}^3 is $\mbox{CHF}_2, \mbox{ CH}_2\mbox{F}, \mbox{ CF}_3 \mbox{ or } \mbox{C}_{1\text{-}4}\mbox{alkyl}.$

Representative examples of R3 include CF3, CH3 and ethyl.

Further representative examples of R3 include CH2F.

In another aspect of the invention R^4 is C_{1-8} alkyl, such as C_{1-3} alkyl.

Representative examples of R4 include CH3.

5 In another aspect of the invention R⁴ is NH₂.

Further representative examples of R4 include NH2.

In another aspect of the invention R^5 is hydrogen or C_{1-3} alkyl.

Representative examples of R5 include H or CH3.

In another aspect of the invention R5 is CN, halogen or CO2Et.

10 Further representative examples of R⁵ include CN, F, CI, CO₂Et.

In another aspect of the invention R^6 and R^7 are independently selected from H or methyl. In another aspect R^6 and R^7 are both H.

In another aspect of the invention A is selected from the group consisting of

where) defines the point of attachment of the ring and A is unsubstituted or substituted by one or two R⁸.

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In another aspect of the invention A is selected from the group consisting of

where defines the point of attachment of the ring

In another aspect of the invention R^8 is selected from the group consisting of halogen, $C_{1:3}$ alkyl, $C_{1:3}$ alkyl substituted by one to three fluorine atoms (e.g. CF_3), and $C_{1:3}$ alkoxy.

5 Representative examples of R⁸ include F, CI, CH₃, methoxy and ethoxy.

Further representative examples of R⁸ include ethyl, fluoromethyl, CF₃ and Br.

Representative examples of B include

In another aspect of the invention R^9 is selected from the group consisting of C_{1-6} alkyl (e.g. ethyl), phenyl and aminomethyl.

In another aspect of the invention R¹⁰ is H.

In another aspect of the invention in compounds of formula (I), (IA) and (IB) n is 0 to 2 (e.g. 1) or in compounds of formula (IC) n is 1 or 2.

In another aspect the invention provides a compound of formula (I) or a pharmaceutically acceptable salt thereof in which:

X is oxvaen:

Y is CH:

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R1 is A(CR6R7)a:

R³ is selected from the group consisting of C₁₋₅alkyl and C₁₋₂alkyl substituted by one to five fluorine atoms:

R4 is C1-salkvl:

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R⁵ is selected from the group consisting of hydrogen, C₁₋₃alkyl, C₁₋₂alkyl substituted by one to five fluorine atoms, C₁₋₃alkylO₂C, halogen, and C₁₋₃alkylS;

A is an unsubstituted 5- or 6-membered heteroaryl or an unsubstituted 6membered aryl, or a 5- or 6-membered heteroaryl or a 6-membered aryl substituted by one or more R8;

R8 is selected from the group consisting of halogen, C1-6alkyl, C1-6alkyl substituted by one more fluorine atoms, C1-salkoxy, and C1-salkoxy substituted by one or more F:

R¹⁰ is selected from the group consisting of H and halogen; and 10 n is 0.

Preferred compounds of the invention are:

4-ethyl-6-[4-(methylsulfonyl)phenyl]-N-(tetrahydro-2H-pyran-4-ylmethyl)-2-

pyridinamine:4-methyl-N-[(1-methyl-1H-pyrazol-4-yl)methyl]-6-[4-15 (methylsulfonyl)phenyl]-2-pyridinamine;

N-[(1,5-dimethyl-1H-pyrazol-4-yl)methyl]-4-methyl-6-[4-(methylsulfonyl)phenyl]-2-pyridinamine:

N-[(1,3-dimethyl-1H-pyrazol-4-yl)methyl]-4-methyl-6-[4-(methylsulfonyl)phenyl]-

2-pyridinamine;

pyridinamine;

4-(6-{[(1,3-dimethyl-1H-pyrazol-4-yl)methyl]amino}-4-ethyl-2-

pyridinyl)benzenesulfonamide;

N-[(1,3-dimethyl-1H-pyrazol-4-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2-pyridinamine;

N-[(1,5-dimethyl-1H-pyrazol-4-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-4-25 (trifluoromethyl)-2-pyridinamine;

4-{4-methyl-6-[(tetrahydro-2H-pyran-4-ylmethyl)amino]-2pyridinyl}benzenesulfonamide:

4-methyl-N-[(1-methyl-1H-pyrazol-3-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-2pyridinamine:

N-(cyclohexylmethyl)-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2pyridinamine;

N-cyclohexyl-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2-pyridinamine;

2-[4-(methylsulfonyl)phenyl]-6-[(2-pyridinylmethyl)oxy]-4-(trifluoromethyl)pyridine; 4-methyl-N-[(3-methyl-4-isoxazolyl)methyl]-6-[4-(methylsulfonyl)phenyl]-2-35

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6-[4-(methylsulfonyl)phenyl]-N-(2-pyridinylmethyl)-4-(trifluoromethyl)-2-pyridinamine;

N-cycloheptyl-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2-pyridinamine; N-(cis-4-methylcyclohexyl)-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2-

5 pyridinamine;

N-(1-ethylpropyl)-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2-pyridinamine; \$N-[(3-methyl-1,2,4-oxadiazol-5-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2-pyridinamine; \$\$

N-[(5-methyl-1,2,4-oxadiazol-3-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-4-10 (trifluoromethyl)-2-pyridinamine;

4-methyl-N-[(1-methyl-1H-pyrazol-5-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-2-pyridinamine;

N-(cyclopentylmethyl)-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2-pyridinamine:

N-[(1-ethyl-1H-1,2,4-triazol-5-yl)methyl]-4-methyl-6-[4-(methylsulfonyl)phenyl]-2-pyridinamine;

4-ethyl-6-[4-(methylsulfonyl)phenyl]-2-[(2-pyridinylmethyl)amino]-3-pyridinecarbonitrile;

4-ethyl-2-[[(5-methyl-2-pyridinyl)methyl]amino}-6-[4-(methylsulfonyl)phenyl]-3-pyridinecarbonitrile:

4-ethyl-2-{[(6-methyl-3-pyridinyl)methyl]amino}-6-[4-(methylsulfonyl)phenyl]-3-pyridinecarbonitrile;

4-ethyl-2-[[(1-methyl-1H-pyrazol-4-yl)methyl]amino}-6-[4-(methylsulfonyl)phenyl]-3-pyridinecarbonitrile;

4-ethyl-6-[4-(methylsulfonyl)phenyl]-2-{[(4-methyl-1,3-thiazol-2-yl)methyl]amino}-3-pyridinecarbonitrile;

4-ethyl-6-[4-(methylsulfonyl)phenyl]-2-[(2-pyridinylmethyl)oxy]-3-pyridinecarbonitrile;

4-ethyl-N-[(1-ethyl-1H-1,2,4-triazol-5-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-2-pyridinamine;

4-ethyl-2-[[(6-methyl-3-pyridinyl)methyl]oxy}-6-[4-(methylsulfonyl)phenyl]-3-pyridinecarbonitrile;

6-[4-(methylsulfonyl)phenyl]-N-[(1-methyl-1H-1,2,4-triazol-5-yl)methyl]-4-(trifluoromethyl)-2-pyridinamine.

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Particularly preferred compounds of the invention are:

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N-cyclohexyl-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2-pyridinamine; 2-[4-(methylsulfonyl)phenyl]-6-[(2-pyridinylmethyl)oxy]-4-(trifluoromethyl)pyridine; 4-methyl-N-[(1-methyl-1H-pyrazol-4-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-2-pyridinamine.

5 It is to be understood that the invention covers all combinations of particular aspects of the invention as described hereinabove.

Since the compounds of the present invention, in particular compounds of formula (I), are intended for use in pharmaceutical compositions, it will be understood that they are each provided in substantially pure form, for example at least 50% pure, more suitably at least 75% pure and preferably at least 95% pure (% are on a wt/wt basis). Impure preparations of the compound of formula (I) may be used for preparing the more pure forms used in pharmaceutical compositions. Although the purity of intermediate compounds of the present invention is less critical, it will be readily understood that the substantially pure form is preferred as for the compounds of formula (I). Preferably, whenever possible, the compounds of the present invention are available in crystalline form

When some of the compounds of this invention are allowed to crystallise or are recrystallised from organic solvents, solvent of recrystallisation may be present in the crystalline product. This invention includes within its scope such solvates. Similarly, some of the compounds of this invention may be crystallised or recrystallised from solvents containing water. In such cases water of hydration may be formed. This invention includes within its scope stoichiometric hydrates as well as compounds containing variable amounts of water that may be produced by processes such as lyophillisation. In addition, different crystallisation conditions may lead to the formation of different polymorphic forms of crystalline products. This invention includes within its scope all the polymorphic forms of the compounds of formula (1).

Compounds of the invention are potent and selective inhibitors of COX-2. This activity is illustrated by their ability to selectively inhibit COX-2 over COX-1.

In view of their selective COX-2 inhibitory activity, the compounds of the present invention are of interest for use in human and veterinary medicine, particularly in the treatment of the pain (both chronic and acute), fever and inflammation of a

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variety of conditions and diseases mediated by selective inhibition of COX-2. Such conditions and diseases are well known in the art and include rheumatic fever; symptoms associated with influenza or other viral infections, such as the common cold; lower back and neck pain; headache; toothache; sprains and strains; myositis; sympathetically maintained pain; synovitis; arthritis, including rheumatoid arthritis; degenerative joint diseases, including osteoarthritis; gout and ankylosing spondylitis; tendinitis; bursitis; skin related conditions, such as psoriasis, eczema, burns and dermatitis; injuries, such as sports injuries and those arising from surgical and dental procedures.

The compounds of the invention are also useful for the treatment of neuropathic pain. Neuropathic pain syndromes can develop following neuronal injury and the resulting pain may persist for months or years, even after the original injury has healed. Neuronal injury may occur in the peripheral nerves, dorsal roots, spinal cord or certain regions in the brain. Neuropathic pain syndromes are traditionally classified according to the disease or event that precipitated them. Neuropathic pain syndromes include: diabetic neuropathy; sciatica; non-specific lower back pain; multiple sclerosis pain; fibromyalgia; HIV-related neuropathy; neuralgia, such as post-herpetic neuralgia and trigeminal neuralgia; and pain resulting from physical trauma, amputation, cancer, toxins or chronic inflammatory conditions. These conditions are difficult to treat and although several drugs are known to have limited efficacy, complete pain control is rarely achieved. The symptoms of neuropathic pain are incredibly heterogeneous and are often described as spontaneous shooting and lancinating pain, or ongoing, burning pain. In addition, there is pain associated with normally non-painful sensations such as "pins and needles" (paraesthesias and dysesthesias), increased sensitivity to touch (hyperesthesia), painful sensation following innocuous stimulation (dynamic, static or thermal allodynia), increased sensitivity to noxious stimuli (thermal, cold, mechanical hyperalgesia), continuing pain sensation after removal of the stimulation (hyperpathia) or an absence of or deficit in selective sensory pathways (hypoalgesia).

The compounds of the invention are also useful for the treatment of other conditions mediated by selective inhibition of COX-2.

For example, the compounds of the invention inhibit cellular and neoplastic transformation and metastatic tumour growth and hence are useful in the

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treatment of certain cancerous diseases, such as colonic cancer and prostate cancer. The compounds of the invention are also useful in reducing the number of adenomatous colorectal polyps and thus reduce the risk of developing colon cancer. The compounds of the invention are also useful in the treatment of cancer associated with overexpression of HER-2/neu, in particular breast cancer.

Compounds of the invention also prevent neuronal injury by inhibiting the generation of neuronal free radicals (and hence oxidative stress) and therefore are of use in the treatment of stroke; epilepsy; and epileptic seizures (including grand mal, petit mal, myoclonic epilepsy and partial seizures).

Compounds of the invention also inhibit prostanoid-induced smooth muscle contraction and hence are of use in the treatment of dysmenorrhoea and premature labour.

Compounds of the invention are also useful in the treatment of liver disease, such as inflammatory liver disease, for example chronic viral hepatitis B, chronic viral hepatitis C, alcoholic liver injury, primary biliary cirrhosis, autoimmune hepatitis, nonalcoholic steatohepatitis and liver transplant rejection.

Compounds of the invention inhibit inflammatory processes and therefore are of use in the treatment of asthma, allergic rhinitis and respiratory distress syndrome; gastrointestinal conditions such as inflammatory bowel disease, Crohn's disease, gastritis, irritable bowel syndrome and ulcerative colitis; and the inflammation in such diseases as vascular disease, migraine, periarteritis nodosa, thyroiditis, aplastic anaemia, Hodgkin's disease, sclerodoma, type I diabetes, myasthenia gravis, multiple sclerosis, sorcoidosis, nephrotic syndrome, Bechet's syndrome, polymyositis, gingivitis, conjunctivitis and myocardial ischemia

Compounds of the invention are also useful in the treatment of ophthalmic diseases such as retinitis, retinopathies, uveitis and of acute injury to the eye tissue.

30 Compounds of the invention are also useful for the treatment of cognitive disorders such as dementia, particularly degenerative dementia (including senile dementia, Alzheimer's disease, Pick's disease, Huntington's chorea, Parkinson's

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disease and Creutzfeldt-Jakob disease), and vascular dementia (including multiinfarct dementia), as well as dementia associated with intracranial space occupying lesions, trauma, infections and related conditions (including HIV infection), metabolism, toxins, anoxia and vitamin deficiency; and mild cognitive impairment associated with ageing, particularly Age Associated Memory Impairment.

Compounds of the invention are also useful in the treatment of disorders ameliorated by a gastroprokinetic agent. Disorders ameliorated by gastroprokinetic agents include ileus, for example post-operative ileus and ileus during sepsis; gastroesophageal reflux disease (GORD, or its synonym GERD); gastroparesis, such as diabetic gastroparesis; and other functional bowel disorders, such as non-ulcerative dyspepsia (NUD) and non-cardiac chest pain (NCCP).

According to a further aspect of the invention, we provide a compound of formula (I) for use in human or veterinary medicine.

According to a further aspect of the invention, we provide a method of treating a human or animal subject suffering from a condition which is mediated by COX-2 which comprises administering to said subject an effective amount of a compound of formula (I).

According to a further aspect of the invention, we provide a method of treating a human or animal subject suffering from an inflammatory disorder, which method comprises administering to said subject an effective amount of a compound of formula (I).

According to another aspect of the invention, we provide the use of a compound of formula (I) for the manufacture of a therapeutic agent for the treatment of a condition which is mediated by COX-2.

According to another aspect of the invention, we provide the use of a compound of formula (I) for the manufacture of a therapeutic agent for the treatment of an inflammatory disorder.

30 It is to be understood that reference to treatment includes both treatment of established symptoms and prophylactic treatment, unless explicitly stated otherwise.

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It will be appreciated that the compounds of the invention may advantageously be used in conjunction with one or more other therapeutic agents. Examples of suitable agents for adjunctive therapy include a 5HT₁ agonist, such as a triptan (e.g. sumatriptan or naratriptan); an adenosine A1 agonist; an EP ligand; an NMDA modulator, such as a glycine antagonist; a sodium channel blocker (e.g. lamotrigine); a substance P antagonist (e.g. an NK1 antagonist); a cannabinoid; acetaminophen or phenacetin: a 5-lipoxygenase inhibitor; a leukotriene receptor antagonist; a DMARD (e.g. methotrexate); gabapentin and related compounds; a tricyclic antidepressant (e.g. amitryptilline); a neurone stabilising antiepileptic drug; a mono-aminergic uptake inhibitor (e.g. venlafaxine); a matrix metalloproteinase inhibitor; a nitric oxide synthase (NOS) inhibitor, such as an iNOS or an nNOS inhibitor; an inhibitor of the release, or action, of tumour necrosis factor α; an antibody therapy, such as a monoclonal antibody therapy; an antiviral agent, such as a nucleoside inhibitor (e.g. lamivudine) or an immune system modulator (e.g. interferon); an opioid analgesic; a local anaesthetic; a stimulant, including caffeine; an H2-antagonist (e.g. ranitidine); a proton pump inhibitor (e.g. omeprazole); an antacid (e.g. aluminium or magnesium hydroxide; an antiflatulent (e.g. simethicone); a decongestant (e.g. phenylephrine, phenylpropanolamine. pseudoephedrine, oxymetazoline. epinephrine. naphazoline, xylometazoline, propylhexedrine, or levo-desoxyephedrine); an antitussive (e.g. codeine, hydrocodone, carmiphen, carbetapentane, or dextramethorphan); a diuretic; or a sedating or non-sedating antihistamine. It is to be understood that the present invention covers the use of a compound of formula (I) in combination with one or more other therapeutic agents.

The compounds of formula (I) are conveniently administered in the form of pharmaceutical compositions. Thus, in another aspect of the invention, we provide a pharmaceutical composition comprising a compound of formula (I) adapted for use in human or veterinary medicine. Such compositions may conveniently be presented for use in conventional manner in admixture with one or more physiologically acceptable carriers or excipients.

As will be appreciated by the person skilled in the art the compounds of the invention may be milled using known milling procedures such as wet milling to obtain a particle size appropriate for tablet formation and for other formulation types. In particular, for those compounds which demonstrate poor bioavailability, finely divided (nanoparticulate) preparations of the compounds of the invention

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may be prepared by processes known in the art, for example see International Patent Application No. WO 02/00196 (SmithKline Beecham).

The compounds of formula (I) may be formulated for administration in any suitable manner. They may, for example, be formulated for topical administration or administration by inhalation or, more preferably, for oral, transdermal or parenteral administration. The pharmaceutical composition may be in a form such that it can effect controlled release of the compounds of formula (I).

For oral administration, the pharmaceutical composition may take the form of, for example, tablets (including sub-lingual tablets), capsules, powders, solutions, syrups or suspensions prepared by conventional means with acceptable excipients.

For transdermal administration, the pharmaceutical composition may be given in the form of a transdermal patch, such as a transdermal iontophoretic patch.

For parenteral administration, the pharmaceutical composition may be given as an injection or a continuous infusion (e.g. intravenously, intravascularly or subcutaneously). The compositions may take such forms as suspensions, solutions or emulsions in oily or aqueous vehicles and may contain formulatory agents such as suspending, stabilising and/or dispersing agents. For administration by injection these may take the form of a unit dose presentation or as a multidose presentation preferably with an added preservative.

Alternatively for parenteral administration the active ingredient may be in powder form for reconstitution with a suitable vehicle.

The compounds of the invention may also be formulated as a depot preparation. Such long acting formulations may be administered by implantation (for example subcutaneously or intramuscularly) or by intramuscular injection. Thus, for example, the compounds of the invention may be formulated with suitable polymeric or hydrophobic materials (for example as an emulsion in an acceptable oil) or ion exchange resins, or as sparingly soluble derivatives, for example, as a sparingly soluble salt.

As stated above, the compounds of the invention may also be used in combination with other therapeutic agents. The invention thus provides, in a

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further aspect, a combination comprising a compound of formula (I) together with a further therapeutic agent.

The combinations referred to above may conveniently be presented for use in the form of a pharmaceutical formulation and thus pharmaceutical formulations comprising a combination as defined above together with a pharmaceutically acceptable carrier or excipient comprise a further aspect of the invention. The individual components of such combinations may be administered either sequentially or simultaneously in separate or combined pharmaceutical formulations.

When a compound of formula (I) is used in combination with a second therapeutic agent active against the same disease state the dose of each compound may differ from that when the compound is used alone. Appropriate doses will be readily appreciated by those skilled in the art.

A proposed daily dosage of a compound of formula (I) for the treatment of man is 0.01mg/kg to 500mg/kg, such as 0.05mg/kg to 100mg/kg, e.g. 0.1mg/kg to 50mg/kg, which may be conveniently administered in 1 to 4 doses. The precise dose employed will depend on the age and condition of the patient and on the route of administration. Thus, for example, a daily dose of 0.25mg/kg to 10mg/kg may be suitable for systemic administration.

Compounds of formula (I) may be prepared by any method known in the art for the preparation of compounds of analogous structure.

Compounds of formula (I) may be prepared by a process which comprises:

reacting a compound R¹XH of formula (II), or a protected derivative thereof, with a compound of formula (III)

$$\mathbb{R}^4 O_2 S$$
 \mathbb{R}^5
 \mathbb{R}^5
 \mathbb{R}^5
 \mathbb{R}^5
 \mathbb{R}^5
 \mathbb{R}^5
 \mathbb{R}^5
 \mathbb{R}^5

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where X is as defined and Z is halogen, such as F, Cl, Br or I, or a sulfonate, such as (4-methyl)benzenesulfonate or trifluoromethanesulfonate and thereafter and if necessary.

interconverting a compound of formula (I) into another compound of formula (I); and/or

deprotecting a protected derivative of compound of formula (I).

The overall synthesis of a compound of formula (I) is shown in Scheme 1 below in which, R^1 to R^3 , R^3 , X and Y are as defined in formula (I) unless otherwise stated, R^4 is C_{1-6} alkyl and Z is a halogen, such as F, Cl, Br or I, or a sulfonate, such as (4-methyl)benzenesulfonate or trifluoromethanesulfonate; LDA is lithium diisopropylamide: THF is tetrahydrofuran.

Referring to Scheme 1, pyridines of formula (I) where R^5 = CI can be obtained by treatment of pyridines of formula (I) where R^5 = H with a chlorinating agent, such as N-chlorosuccinimide, in a solvent, such as acetic acid and at ambient temperature.

Referring to Scheme 1, when X=NR², compounds of formula (I) may be prepared via the treatment of compounds of formula (III) with an amine of formula (III). This is conveniently carried out in a solvent, such as a nitrile (e.g. methylnitrile) and at elevated temperature (e.g. from about 50°C to reflux). An excess of the amine may be used in place of the solvent.

Alternatively, the treatment of compounds of formula (III) with an amine of formula (II) is conveniently carried out in a solvent, such as a tertiary amine (e.g. NMP, N-methyl pyrrolidinone) and at elevated temperature (e.g. from 120°C to 250°C) and with or without microwave irradiation.

Alternatively, the treatment of compounds of formula (III) with an amine of formula (III) may be carried out in the presence of a catalytic quantity of a palladium salt, such as palladium (II) acetate, a phosphine ligand, such as 2,2'-bis(diphenylphosphino)-1,1'-binaphthyl (BINAP), and a base, such as cesium carbonate or sodium tert-butoxide. The reaction is conveniently carried out in a solvent such as toluene or 1,4-dioxan and at elevated temperature.

Alternatively, the treatment of compounds of formula (III) with an amine of formula (II) may be carried out in the presence of a base, such as sodium hydride. The reaction is conveniently carried out in a solvent, such as THF, DMF (N,N-dimethylformamide) or NMP (N-methylpyrrolidinone), at between

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ambient and elevated temperature (e.g. elevated temperature) and with or without microwave irradiation.

Referring to Scheme 1, when X=O, compounds of formula (I) may be prepared by the treatment of compounds of formula (III) with an alcohol of formula (II) in the presence of a base such as sodium hydride. The reaction is conveniently carried out in a solvent such as THF and at between ambient temperature and reflux.

Alternatively, when X=O, compounds of formula (I) may be prepared by treatment of 2-pyridones of formula (IV) with an alkyl halide in the presence of a base , such as silver carbonate, and in a solvent, such as DMF (N,N-dimethylformamide) or n-pentane.

Alternatively, when X=O, 2-pyridones of formula (IV) may be converted to compounds of formula (I) by a Mitsunobu reaction, employing an alcohol of formula (II), a dialkylazodicarboxylate, such as diisopropylazodicarboxylate, a trialkyl- or triarylphosphine, such as tributylphosphine or triphenylphosphine. The reaction is conveniently carried out in a solvent, such as chloroform or THF.

Referring to Scheme 1, 2-pyridones of formula (IV) where $R^5=H$ can be converted to 2-pyridones of formula (IV) where $R^5=F$ by treatment with a fluorinating agent, such as SELECTFLUORTM [1-(chloromethyl)-4-fluoro-1,4,-diazoniabicyclo[2,2,2]octane bis-tetrafluoroborate], in a solvent such as acetonitrile, and at between ambient and elevated temperature (eg elevated temperature)

Referring to Scheme 1, 2-pyridones of formula (IV) where R⁵ = H can be converted to 2-pyridones of formula (IV) where R⁵ = Cl or Br by treatment with a halogenating agent, such as N-chlorosuccinimide or N-bromosuccinimide, in a solvent, such as acetic acid and at ambient temperature.

Referring to Scheme 1, the conversion of 2-pyridones of formula (IV) to the corresponding pyridines of formula (III) where Z is chlorine or bromine, is conveniently carried out employing a phosphorous halide species (e.g. phosphorous (V) chloride) in a solvent, such as a phosphorous oxychalide (e.g. phosphorous oxychloride), and at between ambient and elevated temperature (e.g. elevated temperature). Compounds of formula (III) where Z is chlorine or

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bromine may be converted to compounds of formula (III) where Z is fluorine or iodine using standard interconversion techniques such as those described in 'Comprehensive Organic Transformations: a guide to functional group preparations' by Richard Larock (VCH, 1989), incorporated herein by reference.

- Alternatively, the conversion of 2-pyridones of formula (IV) to the corresponding pyridines of formula (III) where Z is a sulfonate, is conveniently carried out in a solvent, such as a nitrogen-containing solvent (e.g. pyridine) and employing a reagent such as a sulfonyl halide (e.g. (4-methyl)benzenesulfonyl chloride) or a sulfonic anhydride (e.g. tifluoromethanesulfonic anhydride).
- 10 Conveniently the oxidation shown in Scheme 1 is carried out using a monopersulfate compound, such as potassium peroxymonopersulfate (known as Oxone™) and the reaction is carried out in a solvent, such as an aqueous alcohol (e.g. aqueous methanol) and at between -78°C and ambient temperature.
- Alternatively, the oxidation shown in Scheme 1 may be effected using hydrogen peroxide in the presence of sodium tungstate dihydrate. The reaction may be carried out in a solvent such as acetic acid and at between ambient temperature and reflux (e.g. 50°C).
 - Referring to Scheme 1, pyridones of formula (V) are conveniently prepared by treating α , β -unsaturated acids of formula (VII) with two equivalents of LDA in THF at -78°C, followed by a nitrile of formula (VI), according to the procedure described by E. M. Brown, S. Gil, R. Mestres and M. Pavra in *Synthesis*, 2000, 2, pp 273-280, incorporated herein by reference.

Alternatively, pyridones of formulae (IV) and (V) may be prepared as shown in Scheme 2 below.

Referring to Scheme 2, compounds of formula (V) (R^5 = H) may be prepared by treatment of compounds of formula (XIV) with ammonia. The reaction is

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conveniently carried out in a mixture of concentrated aqueous ammonia and dioxane at elevated temperature and in a sealed vessel.

Referring to Scheme 2, compounds of formula (XIV) may be obtained by treatment of compounds of formula (XVII) with a dialkyl malonate (e.g. diethyl malonate) in the presence of a base, such as sodium hydride or a metal alkoxide (e.g. sodium ethoxide). The reaction is conveniently carried out in a solvent, such as THF or an alcohol (e.g. ethanol).

Referring to Scheme 2, compounds of formula (IV) ($\mathbb{R}^5 \neq \mathbb{H}$) may be prepared by treatment of compounds of formula (XV) with a compound of formula (XVI) in the presence of a base, such as sodium hydride or a metal alkoxide (e.g. sodium ethoxide). The reaction is conveniently carried out in a solvent, such as THF or an alcohol (e.g. ethanol).

Referring to Scheme 2 compounds of formula (XIX) may be converted to compounds of formula (XVII) by treatment with an alkynylmetal species, such as an alkynyllithium species or an alkynyl Grignard reagent. The reaction is conveniently carried out in a solvent, such as THF, and at between -78°C and ambient temperature.

Referring to Scheme 2, compounds of formula (XIX) may be obtained by treatment of compounds of formula (XX) with morpholine in the presence of an amide coupling reagent, such as dicyclohexylcarbodiimide (DCC) or 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (EDC) in a solvent such as THF. The reaction may also be carried out in the presence of a base, such as triethylamine or (N,N-diisopropyl)ethylamine.

The synthesis of an intermediate of formula (III) in which R^3 , R^5 and Y are as defined for compounds of formula (I), Z is a halogen, such as F, Cl, Br or I, or a sulfonate such as (4-methyl)benzenesulfonate or trifluoromethanesulfonate, and R^4 is NH₂, is shown in Scheme 3 below. P represents a suitable protecting group.

Referring to Scheme 3, compounds of formula (IX) may be prepared from compounds of formula (VII) in an analogous manner to that described in Scheme 1. Protection of the sulfonamide functionality of the benzonitrile (VIII) may be

achieved using a silicon protecting group, such as the 2-(trimethylsilyl)-ethoxymethyl (SEM) group which can be introduced under standard conditions.

Referring to Scheme 3, the conversion of 2-pyridones of formula (IX) to the corresponding pyridines of formula (III) where Z is halogen, is conveniently carried out employing a phosphorous halide species (e.g. phosphorous (V) chloride) in a solvent, such as a phosphorous oxyhalide (e.g. phosphorous oxychloride), and at between ambient and elevated temperature (e.g. elevated temperature).

Scheme 3

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Alternatively, the conversion of 2-pyridones of formula (IX) to the corresponding pyridines of formula (III) where Z is a sulfonate, is conveniently carried out in a solvent, such as a nitrogen-containing solvent (e.g. pyridine) and employing a reagent such as a sulfonyl halide (e.g. (4-methyl)benzenesulfonyl chloride) or a sulfonic anhydride (e.g. trifluoromethanesulfonic anhydride).

In all alternatives described hereinabove in relation to Scheme 3 for the conversion of (IX) to (III), removal of the protecting groups can be achieved using a source of fluoride, such as tetrabutylammonium fluoride (TBAF), in a

suitable organic solvent such as THF, at a temperature between ambient and reflux.

Conversion of the intermediates of formula (III) to compounds of formula (I) can be achieved as described for Scheme 1. In one variation the nitrogen protecting groups on the sulfonamide functionality may be retained during the transformation of intermediates of formula (III) to compounds of formula (I). In some circumstances removal of the protecting groups occurs during the treatment of intermediate (III) with R¹XH (II). Alternatively, the protecting groups may be removed after treatment of (IIII) with (II) using the standard deprotection conditions described above.

In one variation of Scheme 1, compounds of formula (III) in which Z is halogen, such as F, Cl, Br and I and Y=C, may be synthesised according to Scheme 4 below. R^1 to R^3 , R^5 and Y are as defined in formula (I) unless otherwise stated, R^4 is $C_{4.6}$ alkyl, M represents $B(OH)_2$ or $B(OR)_2$ and m is 0, 1 or 2.

Scheme 4

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Referring to Scheme 4, compounds of formula (XIII) may be converted to compounds of formula (XI) via a Suzuki coupling reaction employing a palladium source, such as palladium tetrakistriphenylphosphine Pd(PPh₃)₄, or Pd₂(dba)₃ and a ligand, such as triphenylphosphine or tri(tertbutyl)phosphine, and a base, such as sodium carbonate, potassium phosphate or potassium fluoride, in a solvent such as a water/ toluene mix, a water/dimethoxyethane mix or 1.4dioxan

Conveniently, the oxidation shown in Scheme 4 is carried out using 3chloroperoxybenzoic acid (m-CPBA) in a chlorinated solvent, such as dichloromethane or chloroform, or a mixture of a chlorinated solvent and aqueous sodium bicarbonate (NaHCO₃). The oxidation is performed at between 0°C and ambient temperature.

Alternatively, the oxidation shown in Scheme 4 may be conveniently carried out in a two-step process, treating compounds of formula (XI) (m = 0) firstly with oxone, and secondly with mCPBA in a chlorinated solvent, such as dichloromethane or chloroform, or a mixture of a chlorinated solvent and aqueous sodium bicarbonate (NaHCO₃). The oxidation is performed at between 0°C and ambient temperature.

The transformation of (X) to the intermediate (III) may conveniently be achieved via treatment of (X) with a phosphorous halide species (e.g. phosphorous (V) chloride) in a solvent, such as a phosphorous oxyhalide (e.g. phosphorous oxychloride), and at between ambient and elevated temperature (e.g. elevated temperature).

Pyridines of formula (XIII) are either known compounds or, when R3 is C1-2alkyl substituted by one to five fluorine atoms, may be prepared from 2chloroisonicotinic acid by standard transformations. For example, when R3 is CHoF or CHFo, this can be conveniently achieved by reduction of 2chloroisonicotinic acid using borane followed by fluorination of the resulting alcohol using a suitable reagent such as DAST, or oxidation of the alcohol, followed by fluorination of the resulting aldehyde with a suitable reagent such as DAST.

It will be appreciated by those skilled in the art that certain of the procedures described in Schemes 1 to 4 for the preparation of compounds of the formula (I)

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or intermediates thereto may not be applicable to some of the possible substituents.

It will be further appreciated by those skilled in the art that it may be necessary to carry out the transformations described in Schemes 1 to 4 in a different order from that described, or to modify one or more of the transformations, to provide the desired compound of formula (I).

It will be appreciated by those skilled in the art that compounds of formula (I) may be prepared by interconversion, utilising other compounds of formula (I) as precursors. Suitable interconversions, such as alkylations, are well known to those skilled in the art and are described in many standard organic chemistry texts, such as 'Advanced Organic Chemistry' by Jerry March, fourth edition (Wiley, 1992), incorporated herein by reference. For example, compounds of other of the compounds of the compound of the compounds of the compound of the compound of the compound of the compound of the corresponding compound of formula (I) wherein R¹ is H.

Acylation of compounds of formula (I) wherein R⁴ is NH₂, to provide compounds of formula (I) wherein R⁴ is R⁹CONH, may be carried out by conventional means, for example by employing conventional acylating agents such as those described in 'Advanced Organic Chemistry', pp 417-424, incorporated herein by reference.

As will be appreciated by those skilled in the art it may be necessary or desirable at any stage in the synthesis of compounds of formula (I) to protect one or more sensitive groups in the molecule so as to prevent undesirable side reactions. The protecting groups used in the preparation of compounds of formula (I) may be used in conventional manner. See, for example, those described in 'Protective Groups in Organic Synthesis' by Theodora W. Greene and Peter G. M. Wuts, third edition, (Wiley, 1999), incorporated herein by reference, which also describes methods for the removal of such groups.

Amines and alcohols of formula (II) are either known compounds or may be prepared by literature methods, such as those described in 'Comprehensive Organic Transformations: a guide to functional group preparations' by Richard Larock (VCH, 1989), incorporated herein by reference.

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Benzonitriles of formula (VI) are either known compounds or may be prepared by literature methods, such as that described by G. Atwell *et al* in *Anti-Cancer Drug Design* 1996, 11, 553, incorporated herein by reference. Where Y = N, nitriles of formula (VI) may be obtained by treating 5-bromo-2-pyridinecarbonitrile with a suitable nucleophile, such as sodium methanethiolate.

 α , β -Unsaturated acids of formula (VII) are either known compounds or may be prepared by literature methods, such as that described by C. Kuroda *et al* in *Tetrahedron* **2000**, *5*6, 6441, incorporated herein by reference.

Certain intermediates described above are novel compounds, and it is to be understood that all novel intermediates herein form further aspects of the present invention. Compounds of formulae (III) and (IV) are key intermediates and represent a particular aspect of the present invention.

Conveniently, compounds of the invention are isolated following work-up in the form of the free base. Pharmaceutically acceptable addition salts of the compounds of the invention may be prepared using conventional means.

Solvates (e.g. hydrates) of a compound of the invention may be formed during the work-up procedure of one of the aforementioned process steps.

The Intermediates and Examples that follow illustrate the invention but do not limit the invention in any way. All temperatures are in °C. Silica chromatography refers to either flash column chromatography performed using Biotage column chromatography cartridges or Solid Phase Extraction (SPE) chromatography, using Varian Mega Bond Elut (Si) cartridges (Anachem) under 15mmHg. Thin layer chromatography (Tlc) was carried out on silica plates. Nuclear magnetic resonance (NMR) spectra were recorded using a Bruker DPX400 spectrometer. Analytical HPLC was conducted on a Supelcosil LCABZ+PLUS column (3.3 cm x 4.6 mm ID) eluting with 0.1% HCO₂H and 0.01 M ammonium acetate in water (solvent A), and 0.05% HCO₂H 5% water in acetonitrile (solvent B), using the following elution gradient 0-0.7 minutes 0%B, 0.7-4.2 minutes linear gradient to 100%B, 4.2-5.3 minutes 0%B, 5.3-5.5 minutes 0%B at a flow rate of 3 ml/minutes. The mass spectra (MS) were recorded on a Waters ZQ mass spectrometer using electrospray positive [(ES+ve to give MH+ and M(NH4)+ molecular ions] or electrospray negative [(ES-ve to give (M-H)-- molecular ion] modes, Mass-directed preparative HPLC was conducted on a Supelco ABZ+

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column ($10 \text{cm} \times 10 \text{mm} \text{ ID}$, $5 \mu \text{m}$) eluting with $0.1 \% \text{ HCO}_2 \text{H}$ in water (solvent A), and $0.05 \% \text{ HCO}_2 \text{H}$ / 5 % water in acetonitrile (solvent B), using the following 10 -minute elution gradients according to the LC retention time: 1.5 - 2.2 mins, 0 - 30 % B; 2.5 - 3.0 mins, 15 - 55 % B; 2.8 - 4.0 mins, 30 - 80 % B; 3.8 - 5.5 mins, 50 - 90 % B. The mass spectra (MS) were recorded on a Micromass ZMD mass spectrometer using electrospray positive [(ES+ve to give MH+ and M(NH4)+ molecular ions] or electrospray negative [(ES-ve to give (M-H)-molecular ion] modes. In addition to those already defined, the following abbreviations are used: Me, methyl; NMP, N- methyl pyrrolidinone; and THF, tetrahydrofuran.

Intermediate 1

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4-Methyl-6-[4-(methylthio)phenyl]-2-pyridone

To a stirred solution of lithium diisopropylamide (50mL of a 2M solution in heptane/THF/ethyl benzene, 0.1mol) in THF (50mL) at -78°C and under an atmosphere of nitrogen was added dropwise a solution of 3-methyl-2-butenoic acid (5g, 0.05mol) in THF (50mL). The reaction was warmed to 0°C for 30 minutes. After cooling to -78°C, a solution of 4-(methylthio)benzonitrile (7.45g, 0.05mol) in THF (50mL) was added dropwise. Upon complete addition, the reaction was warmed to room temperature and stirred for 3 hours. Water (150mL) and ethyl acetate (100mL) were added to the reaction mixture and the resulting precipitate filtered, washed with ethyl acetate and dried to give the title compound (4.96g, 43%) LC retention time 2.75mins, MS m/z 232 (MH⁺).

Intermediate 2

4-Methyl-6-[4-(methylsulfonyl)phenyl]-2-pyridone

To a stirred mixture of intermediate 1 (3.7g, 16.0mmol) in methanol (150mL) at 0°C was added portionwise a suspension of Oxone™ (29.5g, 48.0mmol) in water (100mL). The reaction was warmed to room temperature and stirred for 14 hours. The methanol was removed *in vacuo* and the resulting residue partitioned between saturated aqueous sodium bicarbonate(1L) and chlorofor (500mL) and separated. The aqueous layer was further extracted with chloroform (3 x 200mL) and the combined organic layers were dried over sodium sulfate, filtered and concentrated to give the title compound (3.20g, 76%) LC retention time 2.20mins. MS m/z 264 (MH¹).

Intermediate 3

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4-Methyl-6-[4-(methylsulfonyl)phenyl]pyridine-2-trifluoromethanesulfonate

To a stirred solution of intermediate 2 (3.20g, 12.2mmol) in pyridine (150mL) at 0°C and under an atmosphere of nitrogen was added dropwise trifluoromethanesulfonic anhydride (2.46mL, 14.6mmol). After stirring for 1hr at 0°C, the pyridine was removed in vacuo and the residue partitioned between water (200mL) and dichloromethane (200mL). The layers were separated and the aqueous phase further extracted with dichloromethane (3 x 100mL). The combined organic layers were dried over sodium sulfate, filtered and concentrated in vacuo to give the title compound (4.27g, 89%) LC retention time 3.48mins, MS m/z 396 (MH*).

Example 1

N-Cyclopentyl-4-methyl-6-[4-(methylsulfonyl)phenyl]pyridine-2-amine

A stirred solution of intermediate 3 (60mg, 0.15mmol) and cyclopentylamine (60 μ L, 0.76mmol) in NMP (2mL) was heated at 180°C for 14 hours. Removal of the solvent (vacuum centrifuge) and purification by silica chromatography, eluting with a gradient of cyclohexane to ethyl acetate, gave the title compound (17mg, TLC R_F 0.45, 1:1 ethyl acetate:cyclohexane) MS m/z 331 (MH $^{+}$).

Example 2

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2-Benzyloxy-4-methyl-6-[4-(methylsulfonyl)phenyl]pyridine Route A

To a stirred solution of intermediate 2 (24mg, 0.09mmol) in DMF (0.5mL) was added silver carbonate (28mg, 0.10mmol) followed by benzyl bromide (13 μ L, 0.11mmol). The reaction was stirred at room temperature in the dark for 14h hours before being diluted with diethyl ether (5mL), filtered, washed with water, dried over sodium sulfate, filtered and concentrated *in vacuo* to give the title compound (30mg, 93%) LC retention time 3.54mins, MS m/z 354 (MH*).

Route B

To a stirred suspension of sodium hydride (9mg, 0.22mmol) in DMF (2mL) at room temperature and under an atmosphere of nitrogen was added benzyl alcohol (0.02mL, 0.19mmol). After stirring for 1 hour, the reaction mixture was

added to intermediate 3 (50mg, 0.13mmol) and the reaction heated at 250°C with microwave irradiation. After cooling, the solvent was removed *in vacuo* and the residue partitioned between water (5mL) and dichloromethane (5mL). The layers were separated and the aqueous phase further extracted with dichloromethane (2 x 5mL). The combined organic layers were dried over sodium sulfate, filtered, concentrated *in vacuo* and purified by silica chromatography eluting with a gradient of ethyl acetate in cyclohexane to give the title compound TLC R_F 0.31 (1:3 ethyl acetate:cyclohexane) LC retention time 3.54mins. MS m/z 354 (MH $^+$)

10 Intermediate 4

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2-[4-(methylsulfonyl)phenyl]-4-methylpyridine

To a mixture of 2-chloro-4-methylpyridine (3g, 23.5mmol), 4-(methylsulfonyl)phenylboronic acid (5.64g, 28.2mmol), potassium phosphate (12.0g, 56.4mmol) and DMF (50mL) under an atmosphere of nitrogen was added palladium tetrakistriphenylphosphine (1.36g, 1.18mmol). After heating at 120°C for 14 hours, the reaction was cooled and the DMF removed *in vacuo*. The residue was partitioned between ethyl acetate (100mL) and water (100mL), separated and the organic layer dried over sodium sulfate and concentrated *in vacuo*. Purification by silica chromatography eluting with a gradient of ethyl acetate in cyclohexane gave the title compound (4.29g, 74%) TLC R_F 0.19 (1:1 ethyl acetate:cyclohexane) LC retention time 2.36mins, MS m/z 248 (MH*)

Intermediate 5

2-[4-(methylsulfonyl)phenyl]-4-methylpyridine-N-oxide

A solution of intermediate 4 (3g, 12.2mmol) in dichloromethane (5mL) was added to a solution of 3-chloroperbenzoic acid (7.35g of 57 to 86% grade material) in dichloromethane (15mL) at reflux. After stirring for 3 hours at this temperature, the reaction was cooled, washed sequentially with saturated aqueous sodium bicarbonate solution, saturated aqueous sodium sulfite solution and water, dried over sodium sulfate and concentrated *in vacuo* to give the title compound (3.11g, 97%) LC retention time 1.94mins, MS m/z 264 (MH⁺)

Intermediate 6

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2-Chloro-4-methyl-6-[4-(methylsulfonyl)phenyl]pyridine

A mixture of intermediate 5 (3.11g, 11.8mmol) and phosphorus oxychloride (10mL) was heated at 100°C for 14 hours. After cooling, the reaction was quenched with saturated aqueous sodium bicarbonate solution, with cooling, extracted with dichloromethane and the combined organic extracts dried over sodium sulfate and concentrated *in vacuo*. Purification by silica chromatography eluting with a gradient of ethyl acetate in cyclohexane gave the title compound (1.91g, 58%) TLC R_F 0.35 (1:1 ethyl acetate:cyclohexane) LC retention time 3.13mins. MS m/z 282 (MH⁺)

20 Example 3

N-Benzyl-N-methyl-4-methyl-6-[4-(methylsulfonyl)phenyl]pyridine-2-amine

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A solution of intermediate 6 (10mg, 0.04mmol) and N-methylbenzylamine (20mg, 0.18mmol) in NMP (0.5mL) was heated at 250°C in the microwave for 10minutes. Removal of the solvent (vacuum centrifuge) and purification by silica chromatography, eluting with a gradient of cyclohexane to ethyl acetate, gave the title compound (5mg) LC retention time 3.62mins, MS m/z 367 (MH¹).

Example 83

N-[(1-methyl-1H-pyrazol-4-yl)methyl]-4-methyl-6-[4-(methylsulfonyl)phenyl]pyridine-2-amine

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A stirred solution of intermediate 3 (1.25g, 3.15mmol) and (1-methyl-1H-pyrazol-4-yl)methylamine (0.70g, 6.30mmol) in NMP (10mL) was heated at 180°C for 14 hours, cooled, and loaded evenly onto 5 methanol-conditioned 10g Varian bondelut SCX-2 cartridge. The cartridges were washed with methanol (2 x 40mL each) followed by a solution of 9:1 methanol/concentrated ammonium hydroxide (2 x 40mL each). The ammoniacal fractions were concentrated and purified by silica chromatography eluting with a gradient of cyclohexane to ethyl acetate to give the title compound (780mg) LC retention time 2.32mins, MS m/z 357 (MH*); 14-NMR (CDCl₃) 8 2.23 (3H, s), 3.09 (3H, s), 3.88 (3H, s), 4.47 (2H, d, J = 6Hz), 4.68 (1H, br), 6.28 (1H, s), 6.99 (1H, s), 7.36 (1H, s), 7.50 (1H, s), 8.00 (2H, d, J = 9Hz), 8.19 (2H, d, J = 9Hz).

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1-Ethyl-1H-1,2,4-triazole-5-carbaldehyde

To a solution of 1-ethyl-1*H*-1,2,4-triazole (9.9g, 0.10mol) and N,N,N',N'-tetramethylethylenediamine (15mL) in THF (60mL) at -78°C was added n-butyllithium (64mL of a 1.6M solution in hexanes, 0.10mol). After stirring for 2 hours, DMF (8.7mL, 0.11mol) was added, the reaction allowed to warm to room temperature and stirred for 14 hours before being poured into saturated aqueous sodium bicarbonate solution (300mL). The mixture was extracted with dichloromethane (3 x 150mL) and the combined organics dried over sodium sulfate, filtered and concentrated to give the title compound (>12g) which also contained unreacted starting material ¹H-NMR (CDCl₃) δ 1.48 (3H, t, J = 7Hz), 4.63 (2H, q, 7Hz), 8.03 (1H, s), 10.04 (s, 1H).

1-Ethyl-1H-1,2,4-triazole-5-carbaldehyde oxime

A mixture of crude 1-ethyl-1*H*-1,2,4-triazole-5-carbaldehyde (17.7g), hydroxylamine hydrochloride (12.7g, 0.182mol), sodium bicarbonate (15.3g, 0.182mol) and ethanol (60mL) was heated at reflux for 3 hours. After cooling, the reaction was filtered and the filtrate concentrated *in vacuo*. The resulting residue was crystallised from ethanol to give the title compound (6.17g) ¹H-NMR (d₀-DMSO) 8 1.32 (3H, t, J = 7Hz), 4.41 (2H, q, J = 7Hz), 8.02 (1H, s), 8.25 (1H, s), 12.70 (1H, s)

(1-Ethyl-1H-1,2,4-triazol-5-yl)methylammonium acetate

A mixture of 1-ethyl-1H-1,2,4-triazole-5-carbaldehyde oxime (6.17g, 44mmol), 10% palladium hydroxide on carbon (2.9g) acetic acid (125mL) and ethanol (125mL) were stirred under an atmosphere of hydrogen for 14 hours. The reaction mixture was filtered and concentrated *in vacuo* to give the title compound (7.7g) $^{\rm t}$ H-NMR (d₅-DMSO) δ 1.32 (3H, t, J = 7Hz), 1.89 (3H, s), 3.89 (2H, br), 4.17 (2H, q, J = 7Hz), 7.80 (1H, s).

Example 234

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N-[(1-ethyl-1H-1,2,4-triazol-5-yl)methyl]-4-methyl-6-[4-

10 (methylsulfonyl)phenyl]pyridine-2-amine

(1-ethyl-1H-1,2,4-triazol-5-yl)methylammonium Portions converted to the free base (1-ethyl-1H-1.2.4-triazol-5yl)methylamine by filtering a solution in methanol through an appropriate methanol-conditioned Varian bond-elut aminopropyl cartridge and concentrating the filtrate. A stirred solution of the free base (50mg, 0.40mmol) and intermediate 3 (63mg, 0.16mmol) in NMP (5mL) was heated at 180°C for 14 hours, cooled, and loaded onto a methanol-conditioned 10g Varian bond-elut SCX-2 cartridge. The cartridges were washed with methanol (2 x 40mL) followed by a solution of 9:1 methanol/concentrated ammonium hydroxide (2 x 40mL). The ammoniacal fractions were concentrated and purified by massdirected preparative HPLC to give the title compound (5mg) LC retention time 2.61 mins, MS m/z 372 (MH $^{+}$); ¹H-NMR (CDCI₃) δ 1.42 (3H, t, J = 7Hz), 2.32 (3H, s), 3.10 (3H, s), 4.27 (2H, q, J = 7Hz), 4.84 (2H, d, J = 6Hz), 5.17 (1H, t, J = 6Hz), 6.40 (1H, s), 7.00 (1H, s), 7.85 (1H, s), 8.00 (2H, d, J = 9Hz), 8.13 (2H, d, J = 9Hz).

Intermediate 7

2-[4-(methylthio)phenyl]-4-(trifluoromethyl)-pyridine

To a mixture of 2-chloro-4-(trifluoromethyl)pyridine (19.9g, 0.11mol), 4-(methylthio)phenylboronic acid (21.9g, 0.13mol), 1M aqueous sodium carbonate (180mL) and 1,2-dimethoxyethane (270mL) under an atmosphere of nitrogen was added palladium tetrakistriphenylphosphine (3.78g, 3.3mmol) and the reaction heated at 100°C for 14 hours. After cooling and concentration in vacuo, the residue was partitioned between ethyl acetate (350mL) and water (400mL) and separated. The aqueous layer was further extracted with ethyl acetate (2 x 150mL) and the combined organic layers were dried over sodium sulfate and concentrated in vacuo. Filtration through a pad of silica gel (200g) eluting with a gradient of ethyl acetate in cyclohexane gave the title compound (29.4g) LC retention time 3.62mins, MS m/z 269 (MH¹).

Intermediate 8

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2-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-pyridine

To a stirred suspension of intermediate 7 (29.4g, 0.11mol) in methanol (400mL) at 0°C was added portionwise a suspension of Oxone™ (134g) in water (200mL). The reaction was warmed to room temperature and stirred for 14 hours. The methanol was removed *in vacuo* and the residue diluted with saturated aqueous sodium bicarbonate (2L) and extracted with ethyl acetate (3 x 1L). The combined organic layers were dried over sodium sulfate and

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concentrated in vacuo to give the title compound (32g, 0.106mol) LC retention time 2.90, MS m/z 302 (MH $^{+}$)

Intermediate 9

2-Chloro-4-(trifluoromethyl)-6-[4-(methylsulfonyl)phenyl] pyridine

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To a solution of intermediate 8 (32g, 0.106mol) in dichloromethane (400mL) at reflux was added 3-chloroperbenzoic acid (41.7g of 57 to 86% grade material) portionwise over 15 minutes. After stirring for 14 hours at reflux, the reaction was cooled, diluted with dichloromethane (2L) and washed sequentially with saturated aqueous sodium bicarbonate solution, saturated aqueous sodium sulfite solution containing tetra-n-butylammonium sulfate (4mL) and water, dried sodium sulfate and concentrated in vacuo to (methylsulfonyl)phenyl1-4-(trifluoromethyl)-pyridine-N-oxide (37.2a, containing traces of a tetra-n-butylammonium salt) LC retention time 2.34, MS m/z 318 (MH⁺). A mixture of this crude material and phosphorus oxychloride (110mL) was heated at 110°C for 4 hours. After cooling, the majority of the phosphorus oxychloride was removed in vacuo and the residue neutralised with saturated aqueous sodium bicarbonate solution (300mL), with cooling. The mixture was extracted with chloroform and the combined organic extracts dried over sodium sulfate and concentrated in vacuo. The residue was recrystallised from 2propanol to give the title compound (22.0g) LC retention time 3.23 min, MS m/z 336/338 (MH+).

Example 54

N-cyclohexyl-4-(trifluoromethyl)-6-[4-(methylsulfonyl)phenyl]pyridine-2-amine

A stirred mixture of intermediate 9 (6g, 17.8mmol) and cyclohexylamine (175mL) was heated at 110°C for 14 hours. After cooling, the reaction was diluted with water (1L), acidified with 2N HCI (750mL) and filtered to give the title compound (6.48g) LC retention time 3.81mins MS m/z 399 (MH $^+$); $^+$ H-NMR (CDCl₃) $^-$ 1.22-1.86 (8H, m), 2.60-2.16 (2H, m), 3.09 (3H, s), 3.67-3.78 (1H, m), 4.84 (1H, d, J = 7Hz), 6.57 (1H, s), 7.19 (1H, s), 8.03 (2H, d, J = 9Hz), 8.17 (2H, d, J = 9Hz)

Example 219

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N-(cyclopentanemethyl)- 4-(trifluoromethyl)-6-[4-(methylsulfonyl)phenyl]pyridine-2-amine

A stirred solution of intermediate 9 (630mg, 1.9mmol) and cyclopentanemethylamine (373mg, 3.8mmol) in NMP (5mL) was heated at 180°C for 14 hours. After cooling, the reaction was diluted with water (150mL) and filtered to give the title product (582mg) LC retention time 3.80mins MS m/z 399 (MH⁺); ¹H-NMR (CDCl₃) δ 1.27-1.38 (2H, m), 1.52-1.74 (4H, m), 1.82-1.92 (2H, m) 2.23 (1H, hept, J = 7Hz), 3.10 (3H, s), 3.33 (2H, dd, J = 7Hz & 6Hz), 4.95 (1H, t, J = 6Hz), 6.60 (1H, s), 7.22 (1H, s), 8.03 (2H, d, J = 8Hz), 8.19 (2H, d, J = 8Hz).

Example 208

 $\underline{N-(2-pyridylmethyl)-4-(trifluoromethyl)-6-[4-(methylsulfonyl)phenyl]pyridine-2-amine}$

A solution of intermediate 9 (618mg, 1.84mmol) and 2-pyridylmethylamine (406mg, 3.68mmol) in NMP (4mL) was heated at 250°C with microwave irradiation for 10 minutes. The reaction was diluted with water (100mL) and filtered to give a solid which was further purified by silica chromatography, eluting with a gradient of cyclohexane to ethyl acetate to give the title compound (471mg) LC retention time 2.87mins MS m/z 407 (MH*); ¹H-NMR (CDCl₃) § 3.10 (3H, s), 4.81 (2H, d, J = 5Hz), 6.14 (1H, t, J = 5Hz), 6.76 (1H, s), 7.24 (1H, td, J = 5Hz & 2Hz), 7.37 (1H, d, J = 8Hz), 7.71 (1H, td, J = 8Hz & 2Hz), 8.03 (2H, d, J = 8Hz), 8.19 (2H, d, J = 8Hz), 8.62 (1H, d, J = 5Hz).

Intermediate 10

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4-(Trifluoromethyl)-6-[4-(methylthio)phenyl]-2-pyridone

To a stirred solution of diisopropylamine (11.5mL, 81.8mmol) in THF (75mL) at 0°C was added n-butylithium (51.1mL of a 1.6M solution in hexanes, 81.8mmol). After stirring for 15 minutes, a solution of 4,4,4-trifluoro-3-methyl-2-butenoic acid (6.0g, 38.9mmol) in THF (10mL) was added dropwise. The reaction was allowed to warm to room temperature and stirred for 30 minutes before being cooled to 0°C and treated dropwise with a solution of 4-(methylthio)benzonitrile (2.91g, 19.5mmol) in THF (10mL). Upon complete addition, the reaction was

heated at reflux for 14 hours. After cooling, water (200mL) was added and the mixture extracted with ethyl acetate (250mL). The organic phase was dried over sodium sulfate, filtered and concentrated *in vacuo* and the resulting residue purified by silica chromatography eluting with 1:1 ethyl acetate / cyclohexane to give the title product (2.43g) LC retention time 3.10mins MS m/z 286 (MH⁺).

Intermediate 11

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4-(Trifluoromethyl)-6-[4-(methylsulfonyl)phenyl]-2-pyridone

To a stirred mixture of intermediate 10 (2.43g, 8.52mmol) in methanol (100mL) at 0°C was added portionwise a suspension of Oxone™ (15.7g, 25.6mmol) in water (60mL). The reaction was warmed to room temperature and stirred for 14 hours. The methanol was removed *in vacuo* and the resulting residue partitioned between saturated aqueous sodium bicarbonate(500mL) and chloroform (200mL) and separated. The aqueous layer was further extracted with chloroform (3 x 100mL) and the combined organic layers were dried over sodium sulfate, filtered and concentrated to give the title compound (1.72g) LC retention time 2.57mins, MS m/z 318 (MH*).

Example 164

2-[4-(methylsulfonyl)phenyl]-6-[(2-pyridinylmethyl)oxy]-4-(trifluoromethyl)pyridine

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Diisopropylazodicarboxylate (0.93mL, 4.7mmol) was added dropwise to a solution of intermediate 11 (1g, 3.2mmol), 2-pyridinylmethanol (0.38mL,

3.9mmol) and triphenylphosphine (1.24g, 4.7mmol) in chloroform (80mL). After stirring for 14 hours, the reaction was concentrated and the residue diluted with methanol and loaded onto a methanol-conditioned 10g Varian bond-elut SCX-2 cartridge. The cartridge was washed with methanol (2 x 40mL) followed by a solution of 9:1 methanol/2N hydrochloric acid. The combined acidic fractions were concentrated and the residue triturated with methanol to give the title compound as its hydrochloride salt (348mg) LC retention time 3.35mins, MS m/z 409 (MH*); $^1\text{H-NMR}$ (d₈-DMSO) δ 3.28 (3H, s), 5.79 (2H, s), 7.47 (1H, s), 7.64 (1H, t, J = 6Hz), 7.85 (1H, d, J = 8Hz), 8.03 (2H, d, J = 9Hz), 8.11 (1H, s), 8.17 (1H, t, J = 8Hz), 8.38 (2H, d, J = 9Hz), 8.75 (1H, d, J = 6Hz)

Intermediate 12

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4-{[4-(methylthio)phenyl]carbonyl}morpholine

To a stirred solution of 4-(methylthio)benzoic acid (6.76g, 40.2mmol) and N-[2-(dimethylamino)ethyl]-N-ethylcarbodiimide hydrochloride (9.24g, 48.2mmol) in THF (100mL) was added morpholine (4.2mL, 48.2mmol). After stirring for 2 hours, the reaction was concentrated *in vacuo* and the residue partitioned between ethyl acetate (100mL) and 2M hydrochloric acid (150mL). The organic phase was separated, washed with 1M aqueous sodium carbonate solution, dried over sodium sulfate, filtered and concentrated *in vacuo* to give the title compound LC retention time 3.52mins MS m/z 238 (MH⁺).

Intermediate 13

1-[4-(methylthio)phenyl]-2-pentyn-1-one

To a stirred solution of 1-butyne (approximately 4g) in THF (50mL) at $-78^{\circ}C$ was added dropwise n-butyllithium (47mL of a 1.6M solution in hexanes). Upon

complete addition the reaction was allowed to warm to room temperature and stirred for a further 15 minutes. To the reaction was then added a solution of intermediate 12 (5.97g) in THF (40mL). After stirring for 45 minutes, the reaction was added to a 2:1 mixture of acetic acid and water (150mL) at 0°C. Diethyl ether (50mL) was added and the organic phase separated, washed with saturated aqueous sodium chloride solution, dried over sodium sulfate, filtered and concentrated *in vacuo* to give the title compound (5.09g) LC retention time 3.37mins MS m/z 205 (MH*).

Intermediate 14

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Ethyl 4-ethyl-2-oxo-6-[4-(methylthio)phenyl]-2H-pyran-3-carboxylate

To a stirred solution of sodium ethoxide (0.95g, 13.9mmol) in ethanol (50mL) was added diethyl malonate (10.7mL, 69.4mmol). After stirring for 30 minutes, a solution of intermediate 13 (2.84g, 13.9mmol) in ethanol (50mL) was added and the reaction heated to reflux for 2 hours. After cooling (ice bath), the reaction was acidified to pH ~1 using 2M hydrochloric acid and partitioned between diethyl ether (200mL) and water (50mL). The aqueous phase was further extracted with diethyl ether (2 x 200mL) and the combined organic phases were dried over sodium sulfate, filtered and concentrated *in vacuo*. The resulting crude product was purified by silica chromatography, eluting with a gradient of cyclohexane to ethyl acetate to give the title compound (3.33g) LC retention time 3.47mins MS m/z 319 (MH*).

Intermediate 15

4-Ethyl-6-[4-(methylthio)phenyl]-2(1H)-pyridone

A mixture of intermediate 14 (3.33g, 10.5mmol), concentrated ammonium hydroxide solution (20mL) and 1,4-dioxane (40mL) were heated at 70°C in a sealed vessel for 14 hours. After cooling, the reaction was concentrated to a residue which was triturated with methanol to give the title compound (2.03g) LC retention time 2.87mins MS m/z 246 (MH1°).

Intermediate 16

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4-Ethyl-6-[4-(methylsulfonyl)phenyl]-2(1H)-pyridone

To a stirred mixture of intermediate 15 (2.0g, 8.15mmol) in methanol (60mL) at 0°C was added portionwise a suspension of Oxone™ (15.0g, 24.5mmol) in water (80mL). The reaction was warmed to room temperature and stirred for 14 hours. The methanol was removed *in vacuo* and the resulting residue partitioned between saturated aqueous sodium bicarbonate(100mL) and chloroform (100mL) and separated. The aqueous layer was further extracted with chloroform (3 x 50mL) and the combined organic layers were dried over sodium sulfate, filtered and concentrated to give the title compound (1.98g) LC retention time 2.33mins, MS m/z 278 (MH¹).

Intermediate 17

4-Ethyl-6-[4-(methylsulfonyl)phenyl[pyridine-2-trifluoromethanesulfonate

To a stirred solution of intermediate 16 (1.98g, 7.14mmol) in pyridine (80mL) at 0°C and under an atmosphere of nitrogen was added dropwise trifluoromethanesulfonic anhydride (1.44mL, 8.57mmol), and the reaction was allowed to warm to room temperature. After stirring for 14 hours, the pyridine was removed *in vacuo* and the residue partitioned between water (100mL) and dichloromethane (100mL). The layers were separated and the aqueous phase further extracted with dichloromethane (3 x 50mL). The combined organic layers were dried over sodium sulfate, filtered and concentrated *in vacuo* to give the title compound (2.70g) LC retention time 3.52mins, MS m/z 410 (MH*).

Example 89

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4-Ethyl-6-[4-(methylsulfonyl)phenyl]-N-(tetrahydro-2H-pyran-4-yl)pyridine-2-amine

A stirred solution of intermediate 17 (41mg, 0.10mmol) and tetrahydro-2H-pyran-4-ylamine (21mg, 0.20mmol) in NMP (1mL) was heated at 180°C for 14 hours. After cooling, the reaction was loaded onto a methanol-conditioned 10g Varian bond-elut SCX-2 cartridge. The cartridges were washed with methanol (2 x 40mL) followed by a solution of 9:1 methanol/concentrated ammonium hydroxide (2 x 40mL). The ammoniacal fractions were concentrated and purified by silica chromatography eluting with a gradient of cyclohexane to ethyl acetate to give the title compound (29mg) LC retention time 2.78mins, MS m/z 361 (MH $^{+}$); 1 H-NMR (CDCl₃) δ 1.27 (3H, t, J = 8Hz), 1.57 (2H, qd, J = 11Hz & 4Hz), 2.11 (2H, d,

J = 10Hz), 2.62 (2H, q, J = 8Hz), 3.08 (3H, s), 3.58 (2H, t, J = 10Hz), 3.94 – 4.08 (3H, m), 4.50 (1H, br s), 6.27 (1H, s), 6.96 (1H, s), 7.99 (2H, d, J = 8Hz), 8.14 (2H, d, J = 8Hz)

Intermediate 18

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3-Chloro-4-ethyl-6-[4-(methylsulfonyl)phenyl]-2(1H)-pyridinone

To a stirred solution of intermediate 16 (200mg, 0.72mmol) in acetic acid (5mL) was added N-chlorosuccinimide (96mg, 0.72mmol) and the reaction heated at 90°C for 4 hours. After cooling, the reaction was concentrated *in vacuo* and partitioned between water (25mL) and 4:1 Chloroform/2-propanol (50mL). The organic phase was dried over sodium sulfate, filtered and concentrated *in vacuo* to give the crude title compound (>200mg) LC retention time 2.54mins MS m/z 312/314 (MH*).

Example 220

3-Chloro-4-ethyl-6-[4-(methylsulfonyl)phenyl]-2-[(2-pyridinylmethyl)oxylpyridine

Diisopropylazodicarboxylate (0.076mL, 0.39mmol) was added dropwise to a solution of intermediate 18 (80mg, 0.26mmol), 2-pyridinylmethanol (0.031mL, 0.32mmol) and triphenylphosphine (101mg, 0.39mmol) in chloroform (4mL) After stirring for 14 hours, the reaction was concentrated and the residue diluted with methanol and loaded onto a methanol-conditioned 10g Varian bond-elut

SCX-2 cartridge. The cartridge was washed with methanol (2 x 40mL) followed by a solution of 9:1 methanol/concentrated ammonium hydroxide (2 x 40mL). The ammoniacal fractions were concentrated and purified by mass-directed preparative HPLC to give the title compound (41mg) LC retention time 3.35mins MS m/z 403/405 (MH+*); 1 H-NMR (CDCl₃) δ 1.32 (3H, t, J = 8Hz), 2.87 (2H, q, J = 8Hz), 3.09 (3H, s), 5.70 (2H, s), 7.24 (1H, dd, J = 7Hz & 5Hz), 7.36 (1H, s), 7.60 (1H, d, J = 8Hz), 7.74 (1H, td, J = 8Hz & 2Hz), 7.99 (2H, d, J = 8Hz), 8.14 (2H, d, J = 8Hz), 8.63 (1H, d, J = 5Hz).

Intermediate 19

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1-[4-(Methylsulfonyl)phenyl]-2-pentyn-1-one

To a stirred mixture of intermediate 13 (2.0g, 9.79mmol) in acetonitrile (75mL) at 0°C was added portionwise a suspension of Oxone™ (13.2g, 21.5mmol) in water (75mL). The reaction was warmed to room temperature and stirred for 14 hours. The methanol was removed *in vacuo* and the resulting residue partitioned between water(100mL) and ethyl acetate (100mL) and separated. The organic phase was dried over sodium sulfate, filtered and concentrated to give the title compound (2.24g) LC retention time 2.76mins, MS m/z 237 (MH¹).

Intermediate 20

4-Ethyl-6-[4-(methylsulfonyl)phenyl]-2-oxo-1,2-dihydro-3-pyridinecarbonitrile

To a stirred solution of sodium ethoxide (645mg, 9.5mmol) in ethanol (40mL) was added cyanoacetamide (1.59g, 19.0mmol). After stirring for 15 minutes, a

solution of intermediate 19 (2.24g, 9.5mmol) in ethanol (20mL) was added. Stirring was continued for a further 5 hours, at which time the reaction was made acidic with 2M hydrochloric acid. Water (100mL) was added and the suspension filtered to give the title compound (1.54g) LC retention time 2.42mins MS m/z 303 (MH*).

Example 236

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4-Ethyl-2-{[(6-methyl-3-pyridinyl)methyl]oxy}-6-[4-(methylsulfonyl)phenyl]-3-pyridinecarbonitrile

Diisopropylazodicarboxylate (0.049mL, 0.25mmol) was added dropwise to a solution of intermediate 20 (50mg, 0.17mmol), (6-methyl-3-pyridinyl)methanol (0.023mL, 0.21mmol) and triphenylphosphine (65mg, 0.25mmol) in chloroform (2mL). After stirring for 14 hours, the reaction was diluted with chloroform (10mL), washed with water (10mL), concentrated and the residue triturated with diethyl ether to give the title compound (35mg) LC retention time 2.81mins MS m/z 408 (MH¹); ¹H-NMR (d₀-DMSO) δ 1.29 (3H, t, J = 8Hz), 2.46 (3H, s), 2.85 (2H, q, J = 8Hz), 3.30 (3H, s), 5.64 (2H, s), 7.31 (1H, d, J = 8Hz), 7.84 (1H, dd, J = 8Hz & 2Hz), 7.92 (1H, s), 8.08 (2H, d, J = 8Hz), 8.45 (2H, d, J = 8Hz), 8.63 (1H, d, 2Hz).

20 Intermediate 21

<u>3-Cyano-4-ethyl-6-[4-(methylsulfonyl)phenyl]-2-pyridinyl trifluoromethanesulfonate</u>

To a stirred solution of intermediate 20 (845mg, 2.79mmol) in pyridine (10mL) at 0°C and under an atmosphere of nitrogen was added dropwise trifluoromethanesulfonic anhydride (0.71mL, 4.19mmol), and the reaction was allowed to warm to room temperature. After stirring for 14 hours, the pyridine was removed *in vacuo* and the residue partitioned between water (100mL) and dichloromethane (100mL). The layers were separated and the aqueous phase further extracted with dichloromethane (3 x 50mL). The combined organic layers were dried over sodium sulfate, filtered and concentrated *in vacuo* and the resulting residue purified by silica chromatography eluting with a gradient of cyclohexane to ethyl acetate to give the title compound (1.10g) LC retention time 3.54mins. MS m/z 435 (MH¹).

Example 222

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4-Ethyl-6-[4-(methylsulfonyl)phenyl]-2-[(2-pyridinylmethyl)amino]-3-

15 pvridinecarbonitrile

A stirred solution of intermediate 21 (80mg, 0.18mmol) and 2-pyridinylmethylamine (0.038mL, 0.37mmol) in NMP (1mL) was stirred at room temperature for 14 hours. The reaction was filtered through a methanol-conditioned 5g Varian bond-elut aminopropyl cartridge onto a methanol-conditioned 5g Varian bond-elut SCX-2 cartridge. The SCX-2 cartridge was washed with methanol (2 x 20mL) followed by a solution of 9:1 methanol/concentrated ammonium hydroxide (2 x 20mL). The ammoniacal

fractions were concentrated and the residue triturated with diethyl ether to give the title compound (25mg) LC retention time 2.83mins, MS m/z 393 (MH"); $^1\text{H-}$ NMR (d₆-DMSO) δ 1.27 (3H, t, J = 8Hz), 2.76 (2H, q, J = 8Hz), 3.24 (3H, s), 4.77 (2H, d, J = 6Hz), 7.24 (1H, dd, J = 7Hz & 5Hz), 7.35 (1H, d, J = 8Hz), 7.37 (1H, s), 7.73 (1H, td, J = 8Hz & 2Hz), 7.85 (1H, t, J = 5Hz), 7.95 (2H, d, J = 9Hz), 8.15 (2H, d, J = 9Hz), 8.55 (1H, d, J = 5Hz).

Examples 4 to 236

 Examples 4 to 236, as shown in Tables 1 to 5 that follow, were prepared in the manner described for Examples 1 to 3, 83, 234, 54, 219, 208, 164, 89, 220, 236 and 222 as appropriate.

Table 1

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(I)

R¹ \mathbb{R}^3 R⁵ Ex Х Υ MS 4 4-chlorobenzyl NH CH₃ Н С MH+ 387 5 benzyl NCH₃ CF₃ Н С MH+ 421 С 6 2-furylmethyl NH CF₃ Н MH+ 397 7 C benzvl NH CH₃ н MH+ 353 8 cyclohexanemethyl NH CF₃ н C MH+ 413 9 4-methoxyphenyl NH CH₃ Н C MH+ 369 10 Ω CH₃ н C MH+ 320 2-methylpropyl 0 11 3-pyridyl CH₃ н C MH+ 341 12 allyl NH CF₃ Н С MH+ 357 13 2-chlorophenyl NH CH₃ Н С MH+ 373 14 3,5-difluorobenzyl NH CH₃ Н C MH+ 389 15 NH CH₂ Н С MH+ 354 3-pyridinemethyl

Table 1

 R^1 Εx \mathbb{R}^3 х R⁵ Υ MS 16 4-methoxyphenyl NH CF₃ Н С MH+ 423 17 cyclohexyl NΗ CH₃ Н С MH+ 345 18 n-butvl NH CF₃ Н С MH+ 373 19 2-methylpropyl NH CF_3 Н С MH+ 373 20 4-methoxybenzyl NH CH₂ Н C MH+ 383 21 4-fluorobenzyl NH CH₃ Н С MH+ 371 2-(5-methylfuryl)methyl 22 NΗ CF₃ Н С MH+ 411 23 n-butyl NΗ CH₃ Н С MH+ 319 24 2-furylmethyl NH CH₃ Н С MH+ 343 25 4-methylbenzyl NH CH₃ Н С MH+ 367 26 cyclopentyl NΗ CF₃ Н С MH+ 385 27 4-pyridinemethyl NΗ CH₃ Н С МН+ 354 28 2-pyridinemethyl NH CH_3 С Н MH+ 354 29 2-(6-methylpyridine)methyl NΗ CH_3 Н С MH+ 382 30 4-ethoxybenzyl NH CH₃ Н С MH+ 397 31 2-methylpropyl NH CH_3 Н С MH+ 319 32 propargyl NH CF₃ Н С MH+ 355 33 cyclohexanemethyl NH CH₃ Н С MH+ 359 34 4-pyranylmethyl NΗ CH₃ Н С MH+ 361 35 2-tetrahydrofurylmethyl NH CH₂ Н С MH+ 347

Table 1

$$\bigcap_{\mathsf{CH}_0\mathsf{O}_2\mathsf{S}} \mathsf{R}^{\mathsf{S}}$$

 \mathbb{R}^3 R^5 Υ MS R¹ х Ėx С MH+ 333 NH CH₃ Н 2,2-dimethylpropyl 36 345 Н С MH+ NH СНз 37 2.2.2-trifluoroethyl С MH+ 333 CH₃ Н 38 n-butyl NCH₃ С MH+ 319 39 NEt CH₃ Н ethyl NH CF₃ Н С MH+ 407 40 benzvl NH Н С MH+ 353 CH₃ 41 4-methylphenyl С MH+ 343 NH CH₃ Н 2-furylmethyl 42 С MH+ 357 NH CH₃ Н 4-fluorophenyl 43 359 CH₃ Н С MH+ NH 44 2-thiophenylmethyl MH+ 381 C₂H₅ Н C 45 benzyl NCH₃ С MH+ 375 4-pyranylmethyl NH C_2H_5 Н 46 NH C_2H_5 Н С MH+ 333 47 2-methylpropyl CF₃ н C MH+ 421 NH 48 4-methylbenzyl С MH+ 421 NH CF₃ Н 2-methylbenzyl 49 Н С MH+ 441 NH CF₃ 50 2-chlorobenzyl С MH+ 423 NΗ CF₃ Н 2-(5-methylpyrazine)methyl 51 421 С MH+ NH CF₃ Н 52 (S)-α-methylbenzyl 421 С MH+ 53 (R)-α-methylbenzyl NH CF₃ Н С 399 cvclohexvl NH CF₃ Н MH+ 54 NH CF₃ Н С MH+ 437 55 4-methoxybenzyl

Table 1

(1) R1 \mathbb{R}^3 Ex х R⁵ Y. MS 56 6-methylpyridin-3-yl NH CH₃ Н C MH+ 354 57 benzyl NΗ Н CH₂ С MH+ 353 58 benzyl NCH₃ CH₃ CH₃ С MH+ 381 59 benzyl NΗ CH₃ CH_3 С MH+ 367 60 2-methylpropyl NH CH₃ CH₃ С MH+ 333 61 benzyl NCH₃ Н Н С MH+ 353 62 benzvl NCH₃ CH₃ Н Ν MH+ 368 63 4-methoxybenzyl NH CH₃ Н Ν MH+ 370 64 2-methoxyethyl NH CH₃ С Н MH+ 321 68 2-(6-methylpyridine)methyl NCH₃ CH₃ Н С MH+ 382 69 2-furylmethyl NΗ C_2H_5 Н С MH+ 357 70 4-methoxyphenyl NH CH₃ Н Ν MH+ 370 71 1-methylethyl NΗ CH₃ Н С MH+ 305 74 1-ethylpropyl NΗ CH_3 Н С MH+ 333 75 benzyl NH Н Н C MH+ 339 76 1H-imidazol-2-ylmethyl NH CH₃ Н С MH+ 343 77 1H-pyrazol-4-vlmethyl NH CH₃ Н С MH+ 343 80 (1-methyl-1H-imidazol-2-С NH CH₃ Н MH+ 357 yl)methyl

Table 1

Ex	R ¹	х	R ³	R ⁵	Υ	MS	6 14
81	(3-methyl-1H-pyrazol-4- yl)methyl	NH	CH ₃	Н	С	МН+	357
82	(1-methyl-1H-pyrazol-3- yl)methyl	NH	CH ₃	Н	С	MH+	357
84	1H-imidazol-2-ylmethyl	NH	C ₂ H ₅	Н	С	MH+	357
85	(3-methyl-1H-pyrazol-5- yl)methyl	0	CH₃	Н	С	MH+	358
86	(1-methyl-1H-pyrazol-5- yl)methyl	0	CH ₃	Н	С	MH+	358
87	(1-methyl-1H-1,2,4-triazol- 5-yl)methyl	NH	CH₃	Н	С	MH+	358
88	(5-methyl-3- isoxazolyl)methyl	0	CH₃	Н	С	MH+	359
92	cyclohexyl	NH	CH₂ F	Н	С	MH+	363
93	benzyl	NH	C ₂ H ₅	Н	С	MH+	367
94	(S)-α-methylbenzyl	NH	CH ₃	Н	С	МН+	367
95	2-methylbenzyl	NH	CH ₃	Н	С	MH+	367
96	benzyl	0	C ₂ H ₅	Н	С	мн+	368
97	benzyl	NCH₃	CH ₃	Н	С	MH+	368
98	(6-methyl-3-pyridyl)methyl	NH	CH ₃	Н	С	MH+	368
99	6-methylpyridin-3-yl	NH	C ₂ H ₅	Н	С	MH+	368

Table 1

$$\bigcap_{\mathsf{CH}_3\mathsf{O}_2\mathsf{S}} \mathsf{R}^{\mathsf{S}}$$

 \mathbb{R}^3 R⁵ Y MS R^1 X Ex MH+ 368 NH C₂H₅ Н С 100 benzvl Н С MH+ 369 O C₂H₅ 101 3-pyridylmethyl С 369 C₂H₅ Н MH+ 103 2-pyrazinylmethyl NH С MH+ 371 104 benzvl NΗ CH₂ Н F С 371 C₂H₅ н MH+ NH 105 4-fluorophenyl 371 NH C₂H₅ Н С MH+ 106 2-(5-methylfuryl)methyl С MH+ 371 (2-methyl-1H-imidazol-4-NH C_2H_5 Н 107 yl)methyl NH C_2H_5 Н С MH+ 371 108 (1-methyl-1H-imidazol-2yI)methyl 371 C₂H₅ Н C MH+ 109 (4-methyl-1H-imidazol-5-NH vI)methyl Н (1-methyl-1H-imidazol-2-NCH₃ CH₃ С MH+ 371 110 yl)methyl С MH+ 371 (4-methyl-1H-imidazol-2-NH C_2H_5 Н 111 vI)methvl С 371 112 (1-ethyl-1H-imidazol-2-NH CH₃ Н MH+ yl)methyl CH_3 Н С MH+ 371 (1,3-dimethyl-1H-pyrazol-4-NH 113 yl)methyl

Table 1

3-2-				(1)					
Ex	R ¹	X	R ³	R ⁵	Υ	MS			
114	(1,5-dimethyl-1H-pyrazol-4- yl)methyl	NH	CH ₃	Н	С	MH+	371		
115	(1-methyl-1H-pyrazol-4- yl)methyl	NH	C ₂ H ₅	Н	С	МН+	371		
116	(1-methyl-1H-pyrazol-5- yl)methyl	0	C ₂ H ₅	Н	С	МН+	372		
120	2-thiophenylmethyl	NH	C ₂ H ₅	Н	С	MH+	373		
121	cyclohexyl	NC ₂ H	CH ₃	Н	С	мн+	373		
123	(3-methyl-5- isothiazolyl)methyl	NH	CH ₃	Н	С	MH+	374		
124	(4-methyl-1,3-thiazol-2- yl)methyl	NH	CH ₃	Н	С	мн+	374		
125	(3-methyl-4- isothiazolyl)methyl	NH	CH₃	Н	С	мн+	374		
126	[1-(fluoromethyl)-1H- pyrazol-4-yl]methyl	NH	CH ₃	Н	С	MH+	375		
128	benzyl	NC ₂ H	CH ₃	Н	С	мн+	381		
129	4-methylbenzyl	NH	C ₂ H ₅	Н	С	MH+	381		
131	(1-methyl-1H-pyrazol-4- yl)methyl	NH	CH ₃	CN	С	МН+	382		

Table 1

$$\mathsf{CH_2O_2S} \overset{\mathsf{R}^3}{\longleftarrow} \mathsf{R}^5$$

 \mathbb{R}^3 R^1 R^5 Ex х Υ MS. 132 2-(6-methylpyridine)methyl NH C₂H₅ Н С MH+ 382 С 133 (2-methyl-3-pyridyl)methyl 0 C_2H_5 Н MH+ 383 134 (6-methyl-3-pyridyl)methyl 0 C_2H_5 С 383 Н MH+ 135 2-(6-methylpyridine)methyl 0 C_2H_5 Н C MH+ 383 137 (1-methyl-1H-imidazol-2-NH C_2H_5 Н С MH+ 385 yl)methyl 138 (1,3-dimethyl-1H-pyrazol-4-NH CH₃ Н С MH+ 385 yl)methyl 139 (1.5-dimethyl-1H-pyrazol-4-NH C₂H₅ Н С MH+ 385 yl)methyl 142 (4-methyl-1.3-thiazol-2-NH н С 388 C₂H₅ MH+ yl)methyl 143 (1-methyl-1H-pyrazol-4-NH C₂H₅ F С MH+ 389 vI)methvI 144 [1-(fluoromethyl)-1H-NH C₂H₅ Н С MH+ 389 pyrazol-4-yl]methyl 147 (1-methyl-1H-pyrazol-4-NH CH₃ CI С MH+ 391/ yl)methyl 393 148 NH C_2H_5 CN С MH+ 392 benzvl (6-methyl-3-pyridyl)methyl 0 149 CH_3 CN С MH+ 394 150 3-pvridvl 0 CF₃ Н С MH+ 395

Table 1

$$\mathsf{CH_{5}O_{2}S} \overset{\mathsf{R}^{3}}{\longleftarrow} \mathsf{Y}$$

 R^1 R^3 R⁵ Ex Х Υ MS. 151 benzyl NH C(C Н С MH+ 395 $H_3)_3$ 152 2-(6-methylpyridine)methyl NCH₃ C₂H₅ Н С MH+ 396 153 1H-imidazol-2-vlmethyl NH CF₃ Н С MH+ 397 154 4-ethoxyphenyl NH С C_2H_5 Н MH+ 397 155 tetrahydro-2H-pyran-4-yl NH CF₃ С Н MH+ 401 158 (6-methyl-3-pyridyl)methyl 0 CH₃ С MH+ 369 Н 160 2-methyl-3-pyridyl NΗ CF₃ Н С MH+ 408 162 6-methyl-2-pyridyl NH С CF₃ Н MH+ 408 163 6-methylpyridin-3-vl NH CF₃ н С MH+ 408 165 2-methyl-3-pyridyl 0 CF₃ С MH+ 409 Н 166 0 С 3-pyridylmethyl CF₃ Н MH+ 409 167 6-methylpyridin-3-vl 0 CF₃ С Н MH+ 409 168 2-pyrazinylmethyl NΗ CF₃ С MH+ 409 Н 169 4-fluorophenyl NΗ CF₃ Н C MH+ 411 170 2-furvlmethyl NCH₃ CF₃ Н С MH+ 411 171 (1-methyl-1H-pyrazol-4-NΗ CF₃ Н С MH+ 411 yl)methyl 172 (1-methyl-1H-pyrazol-4-0 CF₃ С MH+ Н 412 yl)methyl

Table 1

 R^1 Ex Х \mathbb{R}^3 R⁵ Υ MS 173 (1-methyl-1H-1,2,4-triazol-NH CF₃ С Н MH+ 412 5-yl)methyl 174 2-thiophenylmethyl NH CF₃ Н С MH+ 413 175 tetrahydro-2H-pyran-4-NH CF₃ Н С MH+ 415 vlmethyl 177 (6-methyl-3-pyridyl)methyl 0 C₂H₅ Н С MH+ 383 178 NH С 2,6-dimethyl-3-pyridyl CF₃ Н MH+ 422 179 (6-methyl-3-pyridyl)methyl NH CF₃ Н С MH+ 422 180 2-(6-methylpyridine)methyl NH CF₃ Н С MH+ 422 181 С 6-ethyl-2-pyridyl NH CF₃ Н MH+ 422 183 0 С 2,6-dimethyl-3-pyridyl CF₃ Н MH+ 423 184 2-(6-methylpyridine)methyl 0 CF₃ Н С MH+ 423 185 0 (2-methyl-3-pyridyl)methyl CF₃ Н C MH+ 423 186 (6-methyl-3-pyridyl)methyl 0 CF₃ С Н MH+ 423 187 (1,3-dimethyl-1H-pyrazol-4-NH CF₃ Н С MH+ 425 yl)methyl 188 (1,5-dimethyl-1H-pyrazol-4-NH CF₃ Н С MH+ 425 yl)methyl 189 (4-methyl-1,3-thiazol-2-NH CF₃ Н С MH+ 428 yl)methyl 190 (5-chloro-3-pyridyl 0 CF₃ Н С MH+ 429

Table 1

R¹ R^3 R^5 Ex X Υ MS 191 6-chloro-3-pyridazinyl NΗ CF₃ Н С MH+ 429/ 431 192 (6-methyl-3-pyridyl)methyl NH CH₃ CN С MH+ 393 193 benzyl NC2 CF₃ С MH+ 435 Н H5 196 2-carboxyphenyl NΗ CF₃ Н С MH+ 437 197 benzyl NH C₂H₅ CO2 C MH+ 439 C_2H_5 200 (5-bromo-2-pyridyl)methyl 0 CF₃ Н С MH+ 486/ 488 201 (3-bromo-4-pyridyl)methyl Ω CF₃ С Н MH+ 486/ 488 202 (3-methyl-4-NH CH_3 Н С MH+ 358 isoxazolyl)methyl 203 5-pyrimidinylmethyl NH CH₃ Н C MH+ 355 204 (1-ethyl-1H-imidazol-2-NH C₂H₅ Н C MH+ 385 yl)methyl 205 (1-methyl-1H-imidazol-2-NCH₃ CH₃ CN С MH+ 396 yl)methyl 206 cis-4-methylcyclohexyl NH CF₃ Н С MH+ 413 207 trans-4-methylcyclohexyl NH CF₃ С MH+ Н 413 209 cycloheptyl NH CF₃ Н С MH+ 413

Table 1

Ex R^1 х \mathbb{R}^3 R⁵ Υ MS 2-pyridylmethyl NH CH₃ CN С MH+ 379 210 211 1-ethylpropyl NH CF₃ Н С MH+ 387 С 212 cyclobutyl NH CF₃ Н MH+ 371 213 (3-methyl-1,2,4-oxadiazol-NH CF₃ Н С MH+ 413 5-vI)methvl 214 (5-methyl-1,2,4-oxadiazol-NH CF₃ Н С MH+ 413 3-yl)methyl 217 2-pyridylmethyl 0 C_2H_5 CN С MH+ 394 218 (1-methyl-1H-pyrazol-5-NH CH₃ Н С HHW 357 yl)methyl 221 trans-4-(ethoxy)cyclohexyl NH CF₃ Н C MH+ 443 С 223 (5-methyl-2-pyridyl)methyl NH C_2H_5 CN MH+ 407 224 (6-methyl-3-pyridyl)methyl NH C₂H₅ CN С MH+ 407 225 (1-methyl-1H-imidazol-2-NH C₂H₅ CN С MH+ 396 yl)methyl 226 (1-ethyl-1H-imidazol-2-NH CN С MH+ 410 C₂H₅ yl)methyl 227 (1-methyl-1H-imidazol-2-NCH₃ C₂H₅ CN С MH+ 410 yl)methyl 228 (1-methyl-1H-pyrazol-4-NH CN С MH+ 396 C₂H₅ yl)methyl

(l)

(1)

Table 1

Ex	R ¹	Х	R ³	R ⁵	Υ	MS	
229	(4-methyl-1,3-thiazol-2- yl)methyl	NH	C ₂ H ₅	CN	С	мн+	413
230	cyclohexyl	NH	C ₂ H ₅	CN	С	MH+	384
231	cyclohexanemethyl	NH	C ₂ H ₅	CN	С	MH+	398
232	(1-ethyl-1H-1,2,4-triazol-5-yl)methyl	NH	C ₂ H ₅	Н	С	мн+	386
233	(1-methyl-1H-1,2,4-triazol- 5-yl)methyl	NH	C ₂ H ₅	Н	С	MH+	372
235	(1-ethyl-1H-1,2,4-triazol-5-yl)methyl	NH	CF ₃	Н	С	MH+	426

Table 2

 \mathbb{R}^3 R^5 Υ MS Ex Н С мн+ 65 C_2H_5 345 С 66 CH₃ CH₃ МН+ 345

Table 3

$$\mathsf{CH_5O_2S} \overset{\mathsf{R}^5}{\underset{\mathsf{R}^{10}}{\bigvee}} \mathsf{R}^5 \qquad \qquad (I)$$

Ex	R ¹	х	R ³	R ⁵	R ¹⁰	Υ	М	S
67	benzyl	NH	CH ₃	Н	F	С	MH+	371

Table 4

 \mathbb{R}^1 X \mathbb{R}^3 MS Ex R⁵ Y 72 n-butyl NH CH₃ Н С MH+ 320 73 2-methylpropyl NH СН3 Н С MH+ 320 cyclohexyl 78 NH CH₃ Н С MH+ 346 С CH₃ 79 benzyl NH Н MH+ 354 tetrahydro-2H-pyran-4-90 NΗ CH₃ Н С MH+ 362 ylmethyl 91 tetrahydro-2H-pyran-4-yl NH Н С 362 C_2H_5 MH+ (6-methyl-3-pyridyl)methyl 102 NH CH₃ Н С MH+ 369 117 (1,5-dimethyl-1H-pyrazol-4-С NH CH_3 Н MH+ 372 yl)methyl 118 (1,3-dimethyl-1H-pyrazol-4-NH CH_3 Н С MH+ 372 yl)methyl

Table 4

X R^5 Ex R¹ R^3 Y MS 119 (1-methyl-1H-pyrazol-4-NH С C_2H_5 Н MH+ 372 vI)methvI 127 tetrahydro-2H-pyran-4-NH C_2H_5 Н С MH+ 376 ylmethyl 136 (6-methyl-3-pyridyl)methyl NΗ C₂H₅ Н С MH+ 383 140 (1,5-dimethyl-1H-pyrazol-4-NΗ C_2H_5 Н С MH+ 386 yl)methyl 141 (1,3-dimethyl-1H-pyrazol-4-NΗ Н С MH+ 386 C₂H₅ yl)methyl 145 (4-methyl-1,3-thiazol-2-NΗ C₂H₅ Н С MH+ 389 yl)methyl 146 (5-chloro-2-pyridyl)methyl NH CH₃ Н C MH+ 389/ 391 3-chloro-4-methylbenzyl 156 NΗ CH₃ Н С MH+ 402/ 404 157 (5-chloro-2-pyridyl)methyl NH C₂H₅ Н C MH+ 403/ 405 161 benzyl NH CF₃ Н С MH+ 408 3-chloro-4-methylbenzyl С 416/ 176 NH C_2H_5 Н MH+ 418 3,4-dichlorobenzyl 182 NH CH₃ Н С MH+ 422/ 424

(l)

(I)

Table 4

Ex	R ¹	X	R ³	R ⁵	Υ	MS	
194	3,4-dichlorobenzyl	NH	C ₂ H ₅	Н	С	MH+	436/ 438
195	3,5-dichlorobenzyl	NH	C ₂ H ₅	Н	С	MH+	436/ 438
199	4-chloro-3- (trifluoromethyl)benzyl	NH	CH₃	Н	С	[M- H]	456

Table 5

Ex	R ⁵	R ⁸	Y	MS	
130	Н	CH ₃	С	MH+	382

Biological Data

Microsomal Assay

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Inhibitory activity against microsomal h-COX2 was assessed against a microsomal preparation from baculovirus infected SF9 cells. An aliquot of microsomal preparation was thawed slowly on ice and a 1/40,000 dilution prepared from it into the assay buffer (sterile water, degassed with argon containing 100mM HEPES (pH 7.4), 10mM EDTA (pH7.4), 1mM phenol, 1mM

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reduced glutathione, 20mg/ml gelatin and 0.001mM Hematin). Once diluted the enzyme solution was then sonicated for 5 seconds (Branson sonicator, setting 4, 1cm tip) to ensure a homogeneous suspension. 155µl enzyme solution was then added to each well of a 96-well microtitre plate containing either 5µl test compound (40x required test concentration) or 5µl DMSO for controls. Plates were then mixed and incubated at room temperature for 1 hour. Following the incubation period, 40μ l of 0.5μ M arachidonic acid was added to each well to give a final concentration of 0.1μ M. Plates were then mixed and incubated for exactly 10 minutes (room temperature) prior to addition of 25μ l M HCl (hydrochloric acid) to each well to stop the reaction. 25μ l of 1M NaOH (sodium hydroxide) was then added to each well to neutralise the solution prior to determination of PGE2 levels by enzyme immunoassay (EIA).

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The following examples had IC $_{50}$ values for inhibition of COX-2 of 0.5 μ M or less and at least a 100-fold selectivity for COX-2 over COX-1, based on comparison of the respective IC $_{50}$ values.

1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 66, 67, 68, 69, 70, 72, 73, 74, 75, 76, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 98, 99, 100, 101, 102, 103, 104, 105, 108, 109, 110, 112, 113, 114, 115, 116, 119, 120, 122, 123, 124, 125, 126, 127, 128, 129, 130, 131, 133, 134, 135, 137, 138, 139, 140, 141, 142, 143, 144, 145, 146, 147, 148, 149, 150, 151, 153, 154, 157, 158, 160, 161, 162, 163, 164, 165, 166, 167, 168, 169, 170, 171, 173, 174, 175, 177, 178, 180, 182, 183, 184, 185, 186, 187, 188, 189, 190, 191, 192, 193, 197, 200, 201, 202, 204, 205, 206, 207, 208, 209, 210, 211, 212, 213, 214, 215, 216, 217, 218, 219, 220, 221, 222, 223, 224, 228, 229, 231, 232, 233, 234, 235, 236.

CLAIMS

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1. A compound of formula (I)

or a pharmaceutically acceptable salt thereof in which:

X is selected from the group consisting of oxygen or NR²;

Y is selected from the group consisting of CH or nitrogen;

 R^1 is selected from the group consisting of H, $C_{1.6}$ alkyl, $C_{1.2}$ alkyl substituted by one to five fluorine atoms, $C_{1.3}$ alkyl $OC_{1.3}$ alkyl, $C_{3.6}$ alkenyl, $C_{3.6}$ alkynyl, $C_{3.10}$ cycloalkyl $C_{0.6}$ alkyl, $C_{4.7}$ cycloalkyl substituted by $C_{1.3}$ alkyl or $C_{1.3}$ alkoxy, $C_{4.12}$ bridged cycloalkyl, $A(CR^5R^7)_n$ and $B(CR^6R^7)_n$;

R² is selected from the group consisting of H and C₁₋₈alkyl; or

R¹ and R², together with the nitrogen atom to which they are attached form a 4-8 membered saturated heterocyclic ring such as a pyrrolidine, morpholine or piperidine ring, or a 5-membered heteroaryl ring which is unsubstituted or substituted by one R⁸:

 R^3 is selected from the group consisting of $\mathsf{C}_{1\text{-}5}$ alkyl and $\mathsf{C}_{1\text{-}2}$ alkyl substituted by one to five fluorine atoms:

R⁴ is selected from the group consisting of C₁₋₆alkyl, NH₂ and R⁹CONH;

 R^5 is selected from the group consisting of hydrogen, $C_{1:3}$ alkyl, $C_{1:2}$ alkyl substituted by one to five fluorine atoms, $C_{1:3}$ alkyl O_2 C, halogen, cyano, ($C_{1:3}$ alkyl O_2 NCO, $C_{1:3}$ alkyl O_3 NCO, $C_{1:3}$ alkyl O_3 S;

 R^6 and R^7 are independently selected from H or $C_{\text{1-8}} \text{alkyl};$

A is an unsubstituted 5- or 6-membered heteroaryl or an unsubstituted 6-membered aryl, or a 5- or 6-membered heteroaryl or a 6-membered aryl substituted by one or more R^6 ;

 R^8 is selected from the group consisting of halogen, C_{1-8} alkyl, C_{1-8} alkyl, substituted by one more fluorine atoms, C_{1-8} alkoxy, C_{1-8} alkoxy substituted by one or more F, NH₂SO₂ and C_{1-8} alkylSO₂;

B is selected from the group consisting of

defines the point of attachment of the ring:

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 R^9 is selected from the group consisting of H, $C_{1-\hat{e}}$ alkyl, $C_{1-\hat{e}}$ alkyl, and $C_{1-\hat{e}}$ alkyl, $C_{1-\hat{e}}$

 R^{10} is selected from the group consisting of H and halogen; and n is 0 to 4.

2. A compound as claimed in claim 1 of formula (IA)

or a pharmaceutically acceptable salt thereof in which:

X is selected from the group consisting of oxygen or NR²:

Y is selected from the group consisting of CH or nitrogen;

 R^1 is selected from the group consisting of H, $C_{1.9}$ alkyl, $C_{1.2}$ alkyl substituted by one to five fluorine atoms, $C_{1.3}$ alkyl $OC_{1.3}$ alkyl, $C_{3.9}$ alkenyl, $C_{3.9}$ alkyl, $C_{3.9}$ cycloalkyl $C_{0.9}$ alkyl, $C_{4.12}$ bridged cycloalkyl, $A(CR^6R^7)_n$ and $B(CR^6R^7)_n$;

 $\ensuremath{R^2}$ is selected from the group consisting of H and $C_{\text{1-6}} alkyl;$ or

R¹ and R², together with the nitrogen atom to which they are attached form a 4-8 membered saturated heterocyclic ring such as a pyrrolidine, morpholine or piperidine ring;

 R^3 is selected from the group consisting of $\mathsf{C}_{1\text{-}5}$ alkyl and $\mathsf{C}_{1\text{-}2}$ alkyl substituted by one to five fluorine atoms:

R⁴ is selected from the group consisting of C₁₋₆alkyl, NH₂ and R⁹CONH;

 R^5 is selected from the group consisting of hydrogen, $C_{1:3}$ alkyl, $C_{1:2}$ alkyl substituted by one to five fluorine atoms, halogen, cyano, $(C_{1:3}$ alkyl)₂NCO, $C_{1:3}$ alkylS and $C_{1:3}$ alkylO₂S:

R⁶ and R⁷ are independently selected from H or C₁₋₆alkyl;

A is an unsubstituted 5- or 6-membered heteroaryl or an unsubstituted 6membered aryl, or a 5- or 6-membered heteroaryl or a 6-membered aryl substituted by one or more R⁸:

 R^8 is selected from the group consisting of halogen, C_{1-6} alkyl, C_{1-6} alkyl, C_{1-6} alkyl substituted by one more fluorine atoms, C_{1-6} alkoxy, C_{1-6} alkoxy substituted by one or more F, NH₂SO₂ and C_{1-6} alkylSO₂;

B is selected from the group consisting of

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$$+ \bigcirc, + \bigcirc, + \bigcirc, + \bigcirc, + \bigcirc$$

defines the point of attachment of the ring:

 R^9 is selected from the group consisting of H, $C_{1-\theta}alkyl,~C_{1-\theta$

 R^{10} is selected from the group consisting of H and halogen; and n is 0 to 4.

3. A compound as claimed in claim 1 of formula (IC)

or a pharmaceutically acceptable salt thereof in which:

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X is selected from the group consisting of oxygen or NR²:

Y is selected from the group consisting of CH or nitrogen;

 R^1 is selected from the group consisting of H, C_{1-2} alkyl, C_{1-2} alkyl substituted by one to five fluorine atoms, C_{1-3} alkyl OC_{1-3} alkyl, C_{3-6} alkenyl, C_{3-6} alkynyl, C_{3-10} cycloalkyl, C_{4-10} cycloalkyl, C_{4-7} cycloalkyl substituted by C_{1-3} alkyl or C_{1-3} alkoxy, C_{4-12} bridged cycloalkyl, $A(CR^6R^7)_n$, and $B(CR^6R^7)_n$;

R² is selected from the group consisting of H and C₁₋₆alkyl; or

R¹ and R², together with the nitrogen atom to which they are attached form a 4-8 membered saturated heterocyclic ring such as a pyrrolidine, morpholine or piperidine ring, or a 5-membered heteroaryl ring which is unsubstituted or substituted by one R⁶.

 R^3 is selected from the group consisting of C_{1-5} alkyl and C_{1-2} alkyl substituted by one to five fluorine atoms;

R⁴ is selected from the group consisting of C₁₋₆alkyl, NH₂ and R⁹CONH;

 R^5 is selected from the group consisting of hydrogen, $C_{1:3}$ alkyl, $C_{1:2}$ alkyl substituted by one to five fluorine atoms, $C_{1:3}$ alkyl O_2 C, halogen, cyano, ($C_{1:3}$ alkyl O_2 C, $C_{1:3}$ alkyl O_3 C, C_1 C,

R⁶ and R⁷ are independently selected from H or C₁₋₆alkyl;

A is an unsubstituted 5- or 6-membered heteroaryl or an unsubstituted 6-membered aryl, or a 5- or 6-membered heteroaryl or a 6-membered aryl substituted by one or more R⁸;

 R^8 is selected from the group consisting of halogen, C_{1-8} alkyl, C_{1-8} alkyl substituted by one more fluorine atoms, C_{1-8} alkoxy, C_{1-6} alkoxy substituted by one or more F, NH₂SO₂ and C_{1-8} alkylSO₂;

B is selected from the group consisting of

defines the point of attachment of the ring;

 R^9 is selected from the group consisting of H, $C_{1-\theta}alkyl,~C_{1-\theta$

 R^{10} is selected from the group consisting of H and halogen; and n is 1 to 4.

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4. A compound as claimed in claim 1 wherein:

X is oxygen:

Y is CH:

R1 is A(CR6R7).:

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R3 is selected from the group consisting of C1.5alkyl and C1.2alkyl substituted by one to five fluorine atoms;

R4 is C1_salkvl:

R⁵ is selected from the group consisting of hydrogen, C₁₋₃alkyl, C₁₋₂alkyl substituted by one to five fluorine atoms, C1-3alkylO2C, halogen, and C1-3alkyIS;

A is an unsubstituted 5- or 6-membered heteroaryl or an unsubstituted 6membered aryl, or a 5- or 6-membered heteroaryl or a 6-membered aryl substituted by one or more R8;

R⁸ is selected from the group consisting of halogen, C₁₋₆alkyl, C₁₋₆alkyl 15 substituted by one more fluorine atoms, C1-6alkoxy, and C1-6alkoxy substituted by one or more F;

R¹⁰ is selected from the group consisting of H and halogen; and n is 0.

- 5. A compound of formula (I) as described in any of Examples 1 to 236. 20
 - 6. A compound of formula (I) selected from the group consisting of: 4-ethyl-6-[4-(methylsulfonyl)phenyl]-N-(tetrahydro-2H-pyran-4-vlmethyl)-2pyridinamine;4-methyl-N-[(1-methyl-1H-pyrazol-4-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-2-pyridinamine:
- N-[(1,5-dimethyl-1H-pyrazol-4-yl)methyl]-4-methyl-6-[4-25 (methylsulfonyl)phenyl]-2-pyridinamine:

N-[(1,3-dimethyl-1H-pyrazol-4-yl)methyl]-4-methyl-6-[4-(methylsulfonyl)phenyl]-2-pyridinamine:

4-(6-{[(1,3-dimethyl-1H-pyrazol-4-yl)methyl]amino}-4-ethyl-2-

30 pyridinyl)benzenesulfonamide:

> N-[(1,3-dimethyl-1H-pyrazol-4-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2-pyridinamine:

> N-[(1,5-dimethyl-1H-pyrazol-4-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2-pyridinamine:

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- 4-{4-methyl-6-[(tetrahydro-2H-pyran-4-vlmethyl)amino]-2-
- pyridinyl}benzenesulfonamide:
- 4-methyl-N-[(1-methyl-1H-pyrazol-3-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-2-pyridinamine:
- 5 N-(cyclohexylmethyl)-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2pyridinamine:
 - N-cvclohexyl-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2pyridinamine:
 - 2-[4-(methylsulfonyl)phenyl]-6-[(2-pyridinylmethyl)oxy]-4-
- 10 (trifluoromethyl)pyridine:

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- 4-methyl-N-[(3-methyl-4-isoxazolyl)methyl]-6-[4-(methylsulfonyl)phenyl]-2pyridinamine:
- 6-[4-(methylsulfonyl)phenyl]-N-(2-pyridinylmethyl)-4-(trifluoromethyl)-2pyridinamine:
- 15 N-cycloheptyl-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2pyridinamine:
 - N-(cis-4-methylcyclohexyl)-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2-pyridinamine;
 - N-(1-ethylpropyl)-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2-
- pyridinamine: N-[(3-methyl-1,2,4-oxadiazol-5-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2-pyridinamine:
 - N-[(5-methyl-1,2,4-oxadiazol-3-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2-pyridinamine:
- 4-methyl-N-[(1-methyl-1H-pyrazol-5-yl)methyl]-6-[4-(methylsulfonyl)phenyl]-25 2-pyridinamine:
 - N-(cyclopentylmethyl)-6-[4-(methylsulfonyl)phenyl]-4-(trifluoromethyl)-2pyridinamine;
 - N-[(1-ethyl-1H-1,2,4-triazol-5-yl)methyl]-4-methyl-6-[4-
- 30 (methylsulfonyl)phenyl]-2-pyridinamine;
 - 4-ethyl-6-[4-(methylsulfonyl)phenyl]-2-[(2-pyridinylmethyl)amino]-3pyridinecarbonitrile:
 - 4-ethyl-2-{[(5-methyl-2-pyridinyl)methyl]amino}-6-[4-(methylsulfonyl)phenyl]-3-pyridinecarbonitrile:
- 35 4-ethyl-2-{[(6-methyl-3-pyridinyl)methyl]amino}-6-[4-(methylsulfonyl)phenyl]-3-pyridinecarbonitrile:

4-ethyl-2-{[(1-methyl-1H-pyrazol-4-yl)methyl]amino}-6-[4-

(methylsulfonyl)phenyl]-3-pyridinecarbonitrile;

 $\hbox{$4$-ethyl-$6-[$4-(methylsulfonyl)]-$2-{[$(4$-methyl-$1,3$-thiazol-$2-dethyl-$1,3$-thiazol-$2-dethyl-$1,3$-thiazol-$2-dethyl-$1,4$-thiazol-$2-dethyl-$1,4$-thiazol-$2-dethyl-$1,4$-thiazol-$2-dethyl-$1,4$-thiazol-$2-dethyl-$1,4$-thiazol-$2-dethyl-$1,4$-thiazol-$2-dethyl-$1,4$-thiazol-$2-dethyl-$1,4$-thiazol-$2-dethyl-$1,4$-thiazol-$2-dethyl-$1,4$-thiazol-$2-dethyl-$1,4$-thiazol-$2-dethyl-$1,4$-thiazol-$2-dethyl-$

yl)methyl]amino}-3-pyridinecarbonitrile;

5 4-ethyl-6-[4-(methylsulfonyl)phenyl]-2-[(2-pyridinylmethyl)oxy]-3pyridinecarbonitrile;

4-ethyl-N-[(1-ethyl-1H-1,2,4-triazol-5-yl)methyl]-6-[4-

(methylsulfonyl)phenyl]-2-pyridinamine;

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4-ethyl-2-{[(6-methyl-3-pyridinyl)methyl]oxy}-6-[4-(methylsulfonyl)phenyl]-3pyridinecarbonitrile; and

6-[4-(methylsulfonyl)phenyl]-N-[(1-methyl-1H-1,2,4-triazol-5-yl)methyl]-4-(trifluoromethyl)-2-pyridinamine.

A process for the preparation of compounds of formula (I) as defined in any
of claims 1 to 6 which comprises reacting a compound R¹XH of formula (II),
or a protected derivative thereof, with a compound of formula (III)

$$\mathbb{R}^4 O_2 S \longrightarrow \mathbb{Y}$$

where X is as defined and Z is halogen or a sulfonate, and thereafter and if necessary, interconverting a compound of formula (I) into another compound of formula (I), and/or deprotecting a protected derivative of compound of formula (I).

- A pharmaceutical composition comprising a compound of formula (I) as defined in any of claims 1 to 6 in admixture with one or more physiologically acceptable carriers or excipients.
- A compound of formula (I) as defined in any of claims 1 to 6 for use in human or veterinary medicine.
 - A method of treating a human or animal subject suffering from a condition which is mediated by COX-2 which comprises administering to said subject

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an effective amount of a compound of formula (I) as defined in any of claims 1 to 6.

11. A method of treating a human or animal subject suffering from an inflammatory disorder, which method comprises administering to said subject an effective amount of a compound of formula (I) as defined in any of claims 1 to 6

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- The use of a compound of formula (I) as defined in any of claims 1 to 6 for the manufacture of a therapeutic agent for the treatment of a condition which is mediated by COX-2.
- 10 13. The use of a compound of formula (I) as defined in any of claims 1 to 6 for the manufacture of a therapeutic agent for the treatment of an inflammatory disorder.

INTERNATIONAL SEARCH REPORT

Internation Application No.

IIN	TERNATIONAL SEARCH REPOR	'	PCT/EP 03	PICEUON NO R/11065
A. CLASS	FICATION OF SUBJECT MATTER		101/11 03	7 11005
A. CLASSI IPC 7	FICATION OF SUBJECT MATTER C07D213/64 A61K31/4418 A61P29/ C07D405/12 C07D413/12 C07D417			0401/12
	o International Patent Classification (IPC) or to both national classific	cation and IPC		
	SEARCHED			
IPC 7	cumentallon searched (classification system followed by classification CO7D A61K A61P	tion symbols)		
Documenta	lion searched other than minimum documentation to the extent that	such documents are incli	uded in the fields s	earohed
Electronic d	ala base consulted during the international search (name of data be	ase and, where practical	, search terms user	d)
	ternal, WPI Data, BEILSTEIN Data, B			
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT			
Category °	Citation of document, with indication, where appropriate, of the re-	elevant passages		Relevant to claim No.
A	WO 01 58881 A (PAYNE JEREMY JOHN NEIL ANTHONY (GB); NAYLOR ALAN (GLAXO) 16 August 2001 (2001-08-1 claim 1	GB);		1-13
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	er documents are listed in the continuation of box C.	χ Patent family r	nembers are listed	in annex.
"A" docume conside "E" earlier d filing di "L" documer which i citation "O" docume other m "P" documer later th	outment but published on or after the international det which may throw doubte on priority, claiming or a which may throw doubte on priority, claiming or or other special transien (set specifical and the priority of the specifical or relevant to an oral disclosure, use, exhibition or same to published prior to the international illing date but an the priority date claimed	"Y" document of particul cannot be ceneider document is combi ments, such combi in the art. "&" document member of	lar relevance; the cred novel or cannot e step when the do lar relavance; the cred to involve an im- ned with one or mo nation being obvious of the same patent	claimed invention be considered to cument is taken alone claimed invention ventive step when the recother such docu- us to a person skilled family
Date of the a	clual completion of the international search	Date of malling of the	ne International sea	arch report
	December 2003	14/01/20	004	
Name and m	ailing address of the ISA European Patient Office, P.B. 5818 Patentlasn 2 NL – 2280 HV Filjswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-2016	Authorized officer Steendij	jk, M	

INTERNATIONAL SEARCH REPORT

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